Polyesters and polyamides
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1.1 Introduction

Among the polycondensation polymers, the most important and widely used today is polyesters; and equally important is polyamides. Polyesters can be processed by melt spinning, injection and/or blow moulding and film extrusion whereas polyamides can be processed by melt spinning and injection moulding and extrusion. More details on polyamides are given in Chapter 2. Polyesters are widely used as packaging materials such as bottles/containers, film, and so on, and also as a fibre, filament, fabrics in textiles and as a base material for photographic film and recording tape applications. The term ‘polyester’ is used for polymeric materials containing ester groups in the polymeric main chains of macromolecules and not to the ester groups in the side chains of the macromolecules, as in the case of poly(vinyl acetate) (or) poly(methyl methacrylate), etc. The term ‘polyester’ is applicable to products derived from dicarboxylic acids and diols. However, the term is not restricted to derivatives of dicarboxylic acids but also to other types of acids such as phosphonic (or) sulphonic (or) phosphoric acids. This chapter is restricted to polyesters derived from dicarboxylic acids and diols alone. Chemical structures of various dicarboxylic acids and diols that are used to make polyesters or copolyesters are given in Table 1.1. Even though large number of polyesters and copolyesters can be prepared by varying dicarboxylic acids and diols and by varying the content of comonomers, only some of them are commercialised and trade names of these polyesters are listed in Table 1.2. Some of the major polyester manufacturers and their capacity are listed in Table 1.3. Even though polyesters have many useful properties, they are blended or alloyed with other polymers to overcome some of the polyester shortcomings. Table 1.4 shows the list of trade names of some of the commercialised polyester blends or alloys.
Table 1.1 Chemical structure of various dicarboxylic acids/diols

<table>
<thead>
<tr>
<th>Dicarboxylic acid/Diol</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terephthalic acid</td>
<td><img src="image1" alt="Structure" /></td>
</tr>
<tr>
<td>Isophthalic acid</td>
<td><img src="image2" alt="Structure" /></td>
</tr>
<tr>
<td>2,6-Naphthalene dicarboxylic acid</td>
<td><img src="image3" alt="Structure" /></td>
</tr>
<tr>
<td>Lactic acid</td>
<td><img src="image4" alt="Structure" /></td>
</tr>
<tr>
<td>Monoethylene glycol</td>
<td><img src="image5" alt="Structure" /></td>
</tr>
<tr>
<td>1,3-Propane diol</td>
<td><img src="image6" alt="Structure" /></td>
</tr>
<tr>
<td>1,4-Butane diol</td>
<td><img src="image7" alt="Structure" /></td>
</tr>
<tr>
<td>1,4-Cyclohexane dimethanol</td>
<td><img src="image8" alt="Structure" /></td>
</tr>
</tbody>
</table>

1.2 Classification

Polyesters are broadly classified into two types, namely: (i) thermoplastic polyesters and (ii) unsaturated polyesters, which upon curing form highly cross-linked thermosets. Thermoplastic polyesters can be further classified into six types: (i) Linear aromatic polyesters, which are mouldable, film-forming and fibre-forming polyesters; (ii) Elastomers, which are block...
Table 1.2 List of trade names of polyesters

<table>
<thead>
<tr>
<th>Polyester</th>
<th>Trade name</th>
<th>Company</th>
<th>Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET</td>
<td>Amilar</td>
<td>Monsanto Co.</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Blue C</td>
<td>DuPont</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Daiya foil</td>
<td>Glanzstoff AG</td>
<td>Fibre</td>
</tr>
<tr>
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<td>Diolen</td>
<td>Fibre</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Dowlex</td>
<td>American Enka Corp</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Encron</td>
<td>Fiber Industries Inc</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Fiber V</td>
<td>VEB Chemiefaserwerk</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Fortrel</td>
<td>Hoechst</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Hostadur</td>
<td>Eastman Chemical Products, Inc</td>
<td>Fibre</td>
</tr>
<tr>
<td>PET</td>
<td>Hostphan</td>
<td>Kurashiki Rayon KK</td>
<td>Fibre</td>
</tr>
<tr>
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<td>Kuraray</td>
<td>VEB Chemiefaserwerk</td>
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</tr>
<tr>
<td>PET</td>
<td>Lavsan</td>
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<tr>
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<tr>
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<td>Mersilene</td>
<td>USSR Government</td>
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<td>Mylar</td>
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</tr>
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<td>Nitron lavsan</td>
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<td>BASF</td>
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<td>Ashley</td>
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<td>Kodel</td>
<td>Eastman Kodak Co.</td>
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<tr>
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<td>Vestan</td>
<td>Chemische Werke Huls</td>
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Table 1.3 List of polyester manufacturers and capacity

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<thead>
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<th>PET suppliers</th>
<th>Capacity (KMTA)</th>
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<td>Reliance Industries Ltd</td>
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<tr>
<td>Indorama</td>
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<td>South Asia Petrochemicals</td>
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<td>JBF</td>
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<td>Eastman Chemicals</td>
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<td>KoSA</td>
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<td>Wellman</td>
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<tr>
<td>M &amp; G</td>
<td>580</td>
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<td>Nanya</td>
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<td>DuPont</td>
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<td>Rhodiaster</td>
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<td>Kohap</td>
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<td>Shin-Kong</td>
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<td>Far Eastern</td>
<td>120</td>
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</tbody>
</table>
Polyester resins

(iii) Liquid crystalline polyesters (LCP), which are typified by very high strain modulus and strength and low strains to break; (iv) Engineering plastics, which are strong, stiff, tough and highly crystalline; (v) Aliphatic polyesters such as polycaprolactone, polylactides, polyglycolides, which are obtained by copolyesters containing stiff and flexible parts; (vi) Aliphatic polyolefinic copolymers containing stiff and flexible parts; (vii) Aliphatic polyesters containing stiff and flexible parts; (viii) Aliphatic polyesters containing stiff and flexible parts.

Table 1.4 List of trade names of polyester blends/alloys

<table>
<thead>
<tr>
<th>Blend/alloy</th>
<th>Trade name</th>
<th>Company</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET/PBT</td>
<td>Valox 800 series</td>
<td>GE Plastics</td>
</tr>
<tr>
<td></td>
<td>Celanex 5000 series</td>
<td>Hoechst-Celanese</td>
</tr>
<tr>
<td>PET/PC</td>
<td>Makroblend</td>
<td>Mobay / Bayer</td>
</tr>
<tr>
<td>PBT/PC</td>
<td>Xenoy</td>
<td>GE Co.</td>
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<td>Makroblend PR</td>
<td>Mobay</td>
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<tr>
<td></td>
<td>Kobblend</td>
<td>Enichem</td>
</tr>
<tr>
<td>PC/PET (or) PBT</td>
<td>Sabre 1600</td>
<td>Dow Chemicals</td>
</tr>
<tr>
<td></td>
<td>Stapron E</td>
<td>DSM</td>
</tr>
<tr>
<td></td>
<td>Edgetek</td>
<td>Polyone</td>
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<tr>
<td>Polyester/PC</td>
<td>Comtuf</td>
<td>ComAlloy</td>
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<td>Eastalloy</td>
<td>Eastman</td>
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<tr>
<td></td>
<td>Valox</td>
<td>GE Plastics</td>
</tr>
<tr>
<td></td>
<td>Naxaloy</td>
<td>MRC Polymers</td>
</tr>
<tr>
<td>PBT/ABS</td>
<td>Cytra</td>
<td>GE Co.</td>
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<td>Dialoy B</td>
<td>Mitsubishi Rayon</td>
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<td>Macalloy B</td>
<td>Japan Synthetic Rubber</td>
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<td>Amoco</td>
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<td>Unitka</td>
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<tr>
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<td>Enuran</td>
<td>GE</td>
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<td>Amoco</td>
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<td></td>
<td>Elitel</td>
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</tr>
</tbody>
</table>

Notes:
* Bisphenol A polyarylates
TPU – Thermoplastic Urethane
TPE – Thermoplastic Elastomer

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ring-opening polymerization of cyclic monomers or dimers; and (vi) Poly(hydroxyl alkanoates) such as poly(3-hydroxy butyrate) [P(3HB)], poly(3-hydroxy valerate), which are found to be biodegradable and biocompatible and are produced from selected carbon substrates such as glucose, gluconates or methanol by a variety of micro-organisms via a fermentation process. Monsanto’s Biopol® is commercially available PHA based polyester.

Linear aromatic commercial polyesters are classified into two types based on the type of aromatic moiety present in the polyester main chain. They are (i) phthalates and (ii) naphthalates. Polyethylene terephthalate (PET), polytrimethylene terephthalate (PTT), polybutylene terephthalate (PBT) belong to the phthalate group because these polyesters were derived from purified terephthalic acid (PTA) (or) dimethyl terephthalate (DMT). Polyethylene isophthalate (PEI) also belongs to phthalate group and was derived from isophthalic acid (IPA) (or) dimethyl isophthalate (DMI). PEI has a glass transition temperature of 51°C. Some of the PTA and DMT manufacturers are Sabic, Mitsubishi, Indian oil, etc. Similarly, Polyethylene naphthalate (PEN), polytrimethylene naphthalate (PTN) and polybutylene naphthalate (PBN) belong to the naphthalate group as these were derived from 2,6-naphthalene dicarboxylic acid (NDA) (or) dimethyl-2,6-naphthalene dicarboxylate (NDC). Both NDA and NDC are commercially produced by BP (British Petroleum). NDC is being manufactured by Mitsubishi Gas Chemical Co. and Amoco. NDA is being manufactured by Frinton Laboratories, Inc. and Kinbester Co. Ltd. When compared to NDC, the availability of NDA is very much limited. The purity of commercial NDA is usually below 95 per cent and the high purity grade (98 per cent) is obtained from alkaline hydrolysis of NDC followed by acidification, which renders this process commercially not viable. NDA and NDC are more expensive than PTA or DMT. Naphthalate moieties in the polymer chain were found to: increase glass transition temperature; increase continuous use temperature; increase heat deflection temperature; increase barrier to oxygen, carbon dioxide, water vapour and gasoline; increase UV absorption cut-off; and increase stiffness and strength but naphthalates are more expensive than terephthalates due to increased cost of NDC/ NDA.

1.3 History

Since ancient times, natural polyesters have been known to mankind. Shellac, natural polyester, as secreted by the lac insect, was used by the ancient Egyptians for embalming mummies. Berzelius first synthesized polyesters of polybasic acids and polyvalent alcohols (Berzelius, 1847) by reacting tartaric acid and glycerol.
Polyesters were initially developed for coating applications and are commonly known as ‘alkyd resins’. The term ‘alkyd’ was coined by combining the first part of the word ‘alcohol’ and the last part of the word ‘acid’. Alkyd resin was first marketed by the General Electric Co. of the USA, with the trade name of Glyptal®, which has become an alternative name to alkyd resin (Kienle and Hovey, 1929). Glyptals were produced by extended condensation reactions involving glycerol, phthalic anhydride and the fatty acids of various oils such as linseed oil, soyabean oil, dehydrated castor oil and tung oil. The unsaturation present in the fatty acids of the oils undergoes aerial oxidation on ageing and become progressively less soluble or insoluble. The oxidation reaction was catalysed by cobalt or manganese naphthenates, which are commonly known as driers. Other polyhydric alcohols such as MEG, propylene glycol/1,3-propane diol (PDO), pentaerythritol and sorbitol and acids namely maleic anhydride, IPA, terephthalic acid and could be used to vary the useful properties of alkyd resins.

Unsaturated polyesters were later developed as laminating resins of fibre reinforced composites. These resins were made by polycondensation reactions involving varied propositions of saturated acids such as phthalic anhydride, IPA, adipic acid and unsaturated acid namely maleic anhydride and a glycol such as MEG, PDO, DEG, etc. These resins could be cured at room temperature using methyl ethyl ketone peroxide and cobalt (or) copper naphthenate (or) octoate. Alternatively, benzoyl peroxide, N, N-dimethyl aniline could be used.

In the late 1920s, W. H. Carothers made a large number of polyesters ranging in molecular weight between 2500 and 5000 by condensation reactions of dicarboxylic acids and diols. In 1930s, Carothers and Arvin (1929) synthesized polyesters having a molecular weight of ~4000 by reacting octadecanedioic acid with PDO (dicarboxylic acids with 5 per cent excess of diols). Carothers and Hill increased the molecular weight to 12000 (Carothers and Hill, 1932). This polymer was known as 3–16 ω ester based on the number of methylene groups in diol and dicarboxylic acid respectively. Later this polymer was known as 3G18 based on the number of carbon atoms in diol and dicarboxylic acid including the carbonyl carbon atom. Carothers and J. W. Hill obtained higher molecular weight polyester up to 25000 by removing the water liberated in the condensation reactions of the step-wise polymerization. Carothers coined the term ‘Super Polymers’ for the polyesters having molecular weights above 10000 because the properties of these polymers were so different from the polymers having low molecular weight i.e., <10000. Super Polyesters were tough, opaque solids which melted to clear viscous liquids at elevated temperatures. Filaments could be drawn by pulling the molten polymer using a glass rod. After allowing the filaments to cool, these filaments could be drawn to
several times their original length. These filaments became transparent, tough and possessed high tenacity.

1.4 Polymerization methods

With respect to the equipment and the conditions followed, the polymerization step in polyesters is similar to that of polyamides. On comparison, the formation of high molecular weight in polyesters differs to some extent. The chemical equilibrium favours the polyamide formation, whereas the equilibrium is much less favoured in polyester; stoichiometric equivalence is easily achieved by the use of salts in polyamides whereas due to the non-formation of salts, the stoichiometric equivalence is more difficult to achieve in polyesters; and the amide interchange reactions are slow, when compared to polyesters.

1.4.1 Step-growth polycondensation

Polyester is formed by step-growth polycondensation from dicarboxylic acid or its diester and diol. The by-products produced are water (or) methanol depending upon the reactant used namely dicarboxylic acid or its dimethyl ester. A number of catalysts are used for the polycondensation reactions such as compounds on antimony, germanium, titanium, aluminium, etc. Antimony based catalysts are used at a dosage level of 300 ppm whereas titanium based catalysts are used at about 20 ppm dosage level. As a result polyesters prepared from titanium based catalysts contain lesser amounts of residual catalyst. Germanium catalyst is expensive and widely preferred in Japan. Enzyme protease from Bacillus licheniformis can also be used as a catalyst for the transesterification of DMT and diol in the synthesis of aromatic polyesters in anhydrous tetrahydrofuran (THF).

1.4.2 Ring-opening polymerization

Polyesters can be prepared by ring-opening polymerization (ROP) (Jerome and Teyssie, 1989; Penczek and Slomkowski, 1989) using cyclic oligomers and polymerization could be achieved at temperatures well below the melting point of the final polymer. Unlike polycondensation reaction, this method does not require removal of any by-products and high molecular weight of >100000 can be achieved in a matter of minutes. Using this method, PET and PBT can be prepared using the corresponding cyclic oligomers. The limitations of this technique are that one has to synthesize cyclic oligomer using diol and diacid chloride and that a solvent and catalyst are required.
1.4.3 Polyaddition reaction

Alternatively, polyesters can be prepared by polyaddition reaction of diepoxides to diacids (Madec and Marechal, 1985). This reaction is catalysed by amines, quaternary ammonium, antimony trioxide, antimony pentachloride, Ph$_3$As. A comprehensive review is reported elsewhere (Madec and Marechal, 1985). Polyaddition reaction is accompanied by the numerous side reactions, which limits its use, where linear polyesters are required. The important side reactions are opening of epoxy ring by water to give hydroxyl pendant groups, which in turn can react with carboxyl end group (or) epoxy ring. These polyesters are used as composites, blends, laminates and biodegradable polymers.

1.4.4 Recycling

Polyesters can be recycled either by physical (or) chemical methods. In the physical recycling method, post-consumer recycled PET flakes are remelted in an extruder for pelletization into chips or direct melt processing into value-added products. Chemical recycling is versatile because it is capable of removing any additive contamination and any type of PET waste can be used. In chemical recycling method, hydrolysis (Pitat, 1959) or methanolysis (Anon, 1959) or glycolysis (MacDowell, 1965) is used. Hydrolysis is carried out under high pressure water or steam or aqueous sodium hydroxide solution at 180–250°C under pressures of 1.4–2 MPa or strong inorganic acid such as nitric acid, sulphuric acid at 85–150°C. Methanolysis involves depolymerization of PET waste in a large quantity of methanol in the presence of catalyst at high pressure and temperature for 3–5 hours (Vaidya, 1988). The depolymerization of PET in the presence of significant excess of glycols such as MEG at elevated temperatures (180–250°C) is termed glycolysis. Post-consumer recycled PET flakes are depolymerised into either the monomers namely PTA or DEG, MEG or into oligomers. Then these monomers or oligomers are reconverted into PET by polycondensation reaction or to unsaturated polyesters by polyaddition reaction. Recycled PET is used for making coarse denier staple fibres (>3 dpf) for carpets, needle-punched non-wovens, tennis ball covers, etc. and for making hollow fibres or as geotextiles for soil stabilisation, silt fencing, rail-bed stabilisation, etc., hollow fibres for filling, partially oriented yarns, dope dyed fibres or filament yarns and strappings for packaging. The use of recycled PET in producing film and sheet products is limited to those applications where clarity, transparency and colour are not of prime importance.
1.5 Poly(ethylene terephthalate) (PET)

The most commercially important aromatic polyester is poly (ethylene terephthalate) (PET). Among the aromatic polyesters, PET is considered as ‘work horse’. PET is otherwise known as polyethylene glycol terephthalate, ethylene terephthalate polymer, poly (oxy-1, 2-ethanediol-1, 4-phenylene dicarbonyl), terephthalic acid-ethylene glycol polyester. It is also known as 2GT. PET is a white or light cream material, has high heat resistance and chemical stability and is resistant to acids, bases, some solvents, oils and fats. The unit molecular weight of PET is 192 and the chemical structure of PET is given below:

The refractive index of PET is 1.574 @ 25°C. PET is insoluble in water and due to hygroscopic nature; it picks up moisture very fast and needs sufficient drying to minimise the drop in intrinsic viscosity (IV) during processing. The abrasion resistance of PET is 60 mg/1000 cycles. PET stands first among all the man-made fibres and challenges cotton with a steady growth rate of about 5 per cent. PET has higher UV absorption cut-off (313 nm) than polycarbonate (290 nm), poly(methyl methacrylate) (285 nm) and high density polyethylene (225 nm). Neither administration of PET powder nor a single administration of chloroform extracts of PET at a dose of 10 g/kg BW had a toxic effect on rats. USFDA approved the use of PET as components of polyethylene phthalate polymers intended for use in contact with food in accordance with the conditions prescribed in 21 CFR part 177.1630. PET fibres display good resistance towards sunlight and are surpassed only by the acrylics. The resistance to mildew, ageing and abrasion is generally considered excellent.

1.5.1 Polymer production

Like any other polyester, there are many routes to produce PET. The most commonly practised ones are: (i) the dimethyl terephthalate (DMT) route and (ii) the purified terephthalic acid (PTA) route. Raw material purity plays an important part with polymer purity. The inhomogeneous deposits in DMT/PTA should be removed by filtration to a fineness range between 1 and 2 μm. Most of the earlier plants were based on the DMT route, since high purity commercial grade PTA was not available during those days.
Nowadays the PTA route is increasingly followed for the production of PET. The PTA route has many advantages over the DMT route. They are:

1. PTA gives higher production output to about 15 per cent higher, due to the faster rate of esterification and reduced raw material weight to product ratio.
2. Capital investment to construct a polyester fibre plant based on PTA is at least 20 per cent less than one based on DMT route.
3. PTA has higher bulk density than DMT and hence requires less storage space compared with DMT.
4. The PTA process needs less MEG as compared with DMT (1:1.15 vs. 1:2). Thus less MEG is to be recovered during polycondensation resulting in lower investment for MEG recovery and recycle system. This in turn promotes energy conservation.
5. PTA process is safe, as methanol is not evolved during esterification while in DMT process; methanol is produced as a by-product. Methanol is highly inflammable and can cause explosion so the system should have explosion proof equipment, which adds to capital cost in DMT process.
6. Small amount of catalyst is used in PTA process. As a result, a much cleaner polymer is obtained, which gives better colour of polymer, higher spin pack life in spinning and lower breakages in stretching and better clarity to containers/bottles.
7. Higher molecular weight is reached in less time in the PTA process than with the DMT process.

Drawback of PTA process is that handling is difficult since PTA does not melt at suitable temperature, nor it is soluble in solvent whereas DMT melts at 140°C and can be easily pumped (or) transferred in molten form. In the case of the PTA process, the reaction proceeds in heterophase due to its poor solubility in MEG, leading to non-uniformity in the product from batch to batch. Hence, efficient stirring during esterification is required. Further PTA process produces higher amount of DEG, a side-reaction product catalysed by acidic medium, than DMT route.

Formation of PET consists of two main reactions namely: (i) esterification (or) precondensation and (ii) polycondensation. The esterification reaction is conducted in excess of MEG. The first step produces a prepolymer which contains bis(hydroxyethyl terephthalate) (BHET) (or) diethylene glycol terephthalate and short chain oligomers and by-products namely water (or) methanol depending upon the raw material used namely PTA or DMT respectively. In the case of the PTA route, the operating temperature was 240–265°C and pressure was 0.4 MPa during precondensation. In order to expedite the rate of polycondensation, a catalyst such as antimony
acetate, antimony trioxide, germanium dioxide or titanium can normally be used. Catalysts other than antimony are listed in Table 1.5. In addition to catalyst, stabiliser namely phosphoric acid, phosphorus acid, trimethyl phosphate, triethyl phosphate should be used to stabilize metal ions such as manganese, zinc, calcium, etc. and deactivate them when used as transesterification catalyst. The most effective catalysts for the transesterification of DMT with MEG were the acetates of zinc, lead (II), mercury (II) together with cobalt (III) acetylacetonate and antimony trioxide (Yoda, 1970). A detailed review for esterification and transesterification reaction using titanium catalyst has been reported elsewhere (Siling, 1996). Theoretically, under such conditions, all chains should terminate with hydroxyl end groups. However, degradation reaction produces a certain amount of carboxyl end groups, which indicates the extent of degradation that had taken place in the melt.

In the polycondensation reaction, there are two chain growth reactions. They are: (i) polyesterification between chain ends with carboxyl and hydroxyl end groups with elimination of water and (ii) polytransesterification between hydroxylethyl end groups and ester end groups with the elimination of MEG, which predominates in the later stages of the reaction. Polycondensation was carried out in vacuum (0.13 KPa) at 275–290°C. At the end of polycondensation, the molten polymer was quenched in water to obtain strands, which were granulated into chips. IV obtained was in

### Table 1.5 Antimony free catalysts

<table>
<thead>
<tr>
<th>Chemical base</th>
<th>Trade name</th>
<th>Company</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Titanium</td>
<td>Ecocat T</td>
<td>Zimmer AG</td>
<td>(Otto 2002)</td>
</tr>
<tr>
<td>Titanium + Phosphorus</td>
<td>Ecocat B</td>
<td>Polytrade</td>
<td>(Otto 2003)</td>
</tr>
<tr>
<td>Titanium complexes</td>
<td>Vertec C400</td>
<td>Johnson Matthey</td>
<td>(Naylor 1997)</td>
</tr>
<tr>
<td>Titanium - Phosphorus complexes</td>
<td>Vertec AC 310</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tetra-isopropyl</td>
<td>Tyzor TPT</td>
<td>Invista/DuPont</td>
<td>(Duan 2002)</td>
</tr>
<tr>
<td>Tetra-n-butyl</td>
<td>Tyzor nBT</td>
<td></td>
<td>(Lustig and</td>
</tr>
<tr>
<td>titanate</td>
<td></td>
<td></td>
<td>Burch 1999)</td>
</tr>
<tr>
<td>Titanium – Silicon</td>
<td>C94</td>
<td>Accordis</td>
<td>(Martl 1995)</td>
</tr>
<tr>
<td>Alkali titanate</td>
<td>Hombifast</td>
<td>Sachtleben Chemie</td>
<td>(Schmidt 1996)</td>
</tr>
<tr>
<td>Germanium</td>
<td>TG/19</td>
<td>Teck Cominco (formerly Meld Form)</td>
<td>(John 2001)</td>
</tr>
<tr>
<td>Aluminium</td>
<td></td>
<td>Toyobo</td>
<td>(Nakajima 2002)</td>
</tr>
</tbody>
</table>
the range of 0.5 to 0.7 dL/g and the residual acetaldehyde (AA) content was approx. 50 ppm.

Side reactions

Many side reactions take place during polymerization of PET. The important one is the etherification of MEG to form diethylene glycol (DEG). Up to 50 per cent of total DEG is generated during the initial stages of polycondensation in the preheating and another 40 per cent in the low vacuum state and remaining 10 per cent during the final vacuum stage (Renwen, 1983). Being a diol and less volatile, DEG gets incorporated as a comonomer in PET. In the PTA route, DEG is formed during the esterification step. Both alkali and alkaline earth metals as well as quaternary ammonium compounds can be used to bring about a significant reduction in the level of DEG formed. DMT route produces less DEG content than PTA route because high acidity of PTA catalyses the formation of DEG and this must be controlled at an acceptable level to give consistent polymer properties. The effect of increasing DEG content on PET properties leads to: (i) reduction in heat and light resistance; (ii) decrease in the glass transition and melting temperature; (iii) decrease in the hydrolytic thermal and oxidative degradation resistance; (iv) increase in the dyeability up to 1.5 to 2.5 per cent; (v) slowness in the crystallization kinetics; and (vi) increase in the softness and the breakage of pills. For textile applications, it is necessary to ensure that the DEG content is constant from batch to batch. Otherwise, it leads to non-uniform dyeing. The decrease in melting point ($T_m$) with respect to DEG content can be predicted with an accuracy of ±1.5 per cent using the following equation:

$$T_m ({}^\circ\text{C}) = 261 - (\text{wt. } \% \text{ of DEG } \times 4.2)$$

DEG further loses water to form dioxane, which has higher vapour pressure and will be removed from the process as the column top product. Dioxane is formed when the DEG end group undergoes intramolecular reaction to form the terephthalic acid end group and dioxane (Hovenkamp and Munting, 1970).

Thermal degradation occurs during synthesis and processing at temperatures above melting point. As a result, the ester bond undergoes scission, leading to the terminal vinyl group and carboxyl end group, which then rearrange to form the AA. In packaging application, the AA migrates to the food content which affects the flavour of the food. The acceptable limit is <1 ppm. Another side reaction is the formation of cyclic oligomers, which occurs by ester interchange reaction in PET and cyclisation occurs mainly at the hydroxyl chain ends in the molten state (Cho et al., 1998).
1.5.2 Solid state polymerisation

Pilati (1989) has published a comprehensive review on solid state polymerisation (SSP). To further reduce acetaldehyde content and increase molecular weight, the SSP is carried out for the amorphous chips. The advantages of choosing the SSP route rather than the melt phase route for further increasing the molecular weight are as follows:

1. problems associated with the stirring of the viscous melt are eliminated in the solid state;
2. lower investment and running costs of the continuous SSP route, which does not require very high temperatures and vacuum associated with the melt phase; and
3. degradation and side reactions were limited in SSP due to the lower processing temperatures used.

Before carrying out SSP, amorphous chips are crystallised in fluidised pre-crystalliser at about 170°C for 30 min with vigorous agitation to prevent sticking. The chips are further crystallised at 190°C in a rotary crystalliser, giving them a crystallinity of 30 per cent and density of 1.3845 g/cc. Then SSP is carried out by gravity flow continuous SSP reactor. SSP is operated in an inert gas stream or in a vacuum and an operating temperature of 180–240°C. Nitrogen gas is preferably used. IV of PET chips increased to 0.72–1.2 dL/g and the residual AA content was below 1 ppm.

It has been reported that the molecular weight correlates with the square root of the reaction time for both catalysed and uncatalysed SSP process (Jabarín and Lofgren, 1986; Droscher and Wegner, 1978). Rate of SSP reaction depends on many factors such as temperature and time (Jabarín and Lofgren, 1986), pellet size (Chen and Chen, 1987), crystallinity (Chang, 1970), additive types and concentration (Kokkolas et al., 1995), process gas type and quantity (Hsu, 1967; Mallon et al., 1998), molecular weight (Buxbaum, 1979) and end group concentration (Wu et al., 1997). Ravindrath and Mashelkar (1990) developed a method for the typical industrial SSP process of PET. They concluded that the reaction rate decreases by a factor of 6 for the temperature range between 285 and 220°C accompanied by a decrease of the thermal degradation by a factor of 40.

The SSP process can be eliminated, if an integrated process such as the direct high IV (DHI) process (or) melt-to-resin (MTR) technology is followed. DHI and MTR technologies were developed by Zimmer and Uhde Inventa with Fischer respectively. The integrated process gives economic advantage due to reduced investment and conversion cost, reduced raw material consumption and better/comparable chip quality. Table 1.6 shows the comparison of PET resin quality produced by MTR and SSP process.
Polyester resins

1.5.3 Properties

Polyesters have good resistance to most mineral acids but concentrated sulphuric acid dissolves polyesters with partial decomposition. Polyesters display excellent resistance to conventional bleaching agents, cleaning solvents and surfactants and the degree of crystallinity and molecular orientation determines the extent of resistance. Basic substances attack polyester fibres in two ways. Strong alkalis cause dissolution of the fibre surface. Weak bases such as ammonia and other organic bases such as methyl amine penetrate into the non-crystalline regions of the structure. Because of the lack of chemical dye sites, polyester fibres are usually dyed with disperse dyes such as amacron, artisil, calcosperse, cekryl, celliton, dispersol, duranol, esterophile, foron, genecron, harshaw ester, latyl, palanil, polydye, resolin, samaron, setacyl, terasil, etc. Polyester fibres are dyed from an aqueous bath at above 100°C or by the use of a carrier such as biphenyl, phenyl salicylate. The rate of dyeing polyester fibres is slower than that of cellulose triacetate or acetate.

Intrinsic viscosity (IV) is a measure of molecular weight and that determines the end use of PET. Solvents such as o-chlorophenol, 60:40 phenol/tetrachloroethane and 50:50 phenol/1,2-dichlorobenzene are typically used to determine IV of PET resin. IV is correlated to the molecular weight by the well-known empirical relationship proposed by Mark and Houwink:

<table>
<thead>
<tr>
<th>Processes</th>
<th>Unit</th>
<th>MTR°</th>
<th>SSP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resin IV</td>
<td>dL/g</td>
<td>0.65–0.85</td>
<td>0.75–0.82</td>
</tr>
<tr>
<td>IV tolerance</td>
<td></td>
<td>±0.008</td>
<td>±0.02</td>
</tr>
<tr>
<td>ΔIV core/shell</td>
<td></td>
<td>0</td>
<td>&gt;0.03</td>
</tr>
<tr>
<td>Precursor COOH°</td>
<td>meq/kg</td>
<td>50–70</td>
<td>30–40</td>
</tr>
<tr>
<td>COOH°</td>
<td>meq/kg</td>
<td>&lt;25</td>
<td>&lt;30</td>
</tr>
<tr>
<td>Melting point ⊗</td>
<td>°C</td>
<td>248–252</td>
<td>248–252</td>
</tr>
<tr>
<td>Colour b° ⊗ CIE Lab</td>
<td></td>
<td>&lt;+1.0</td>
<td>&lt;+1.0</td>
</tr>
<tr>
<td>Crystallinity</td>
<td>%</td>
<td>&lt;35</td>
<td>&gt;50</td>
</tr>
<tr>
<td>Heat of fusion</td>
<td>J/g</td>
<td>42–48</td>
<td>&gt;50</td>
</tr>
<tr>
<td>Polydispersity</td>
<td></td>
<td>1.8</td>
<td>2.4</td>
</tr>
</tbody>
</table>

*Note:*

°Depends on the recipe

Uhde Inventa Fischer claims that MTR can render a benefit of €30 to €40 on conversion costs compared with older conventional continuous plants.
\[ [\eta] = K \times \bar{M}^\alpha \]

where \( \eta \) is IV, \( \bar{M} \) is the molecular weight and \( K \) and \( \alpha \) are constants, which are solvent dependent. Depending upon how the co-relationship was established \( M \) can be expressed as, number average molecular weight (\( M_n \)), weight average molecular weight (\( M_w \)) and viscosity average molecular weight (\( M_v \)).

Ravens and Ward (Ravens and Ward, 1961) have established the above relationship using o-chlorophenol at 25°C by end-group analysis.

\[ [\eta] = 1.7 \times 10^{-4} \times M_n^{0.83} \]

Melt viscosity (low shear) of PET can be predicted from the IV, which was measured using o-chlorophenol, using the formula

\[ \eta_0 = 0.129 \times [\eta]^{5.35} \exp\{6800/T\} \]

where \( \eta_0 \) is the Newtonian melt viscosity (poise) and \( T \) is the absolute temperature.

When GPC was used the above relationship changed as follows:

\[ [\eta] = 1.47 \times 10^{-4} \times M_w^{0.768} \]
\[ [\eta] = 4.68 \times 10^{-4} \times M_w^{0.68} \]

In the above relationships, IV solvents used were o-chlorophenol and 60:40 phenol/tetrachloroethane at 25°C (Moore, 1960) respectively.

IV values of 0.40, 0.63, 0.72 and 1.0 dL/g correspond to number of average molecular weights of approx. 10000, 18000, 24000 and 40000 g/mol respectively. IV of fibre grade is in the range of 0.40 to 0.70 dL/g and yarns in the range of 0.72 to 0.98 dL/g. Recommended IV of various fibre types are given in Table 1.7. IV of bottle grade is in the range of 0.70 to 0.85 dL/g. IV of biaxially oriented film and thermoforming sheet grades is in the range of 0.60 to 0.70 dL/g and 0.7 to 1.0 dL/g respectively. IV of

<table>
<thead>
<tr>
<th>Types of fibre/yarn</th>
<th>IV (dL/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low pill staple fibre</td>
<td>0.40–0.50</td>
</tr>
<tr>
<td>Wool type fibre</td>
<td>0.58–0.63</td>
</tr>
<tr>
<td>Cotton type fibre</td>
<td>0.58–0.64</td>
</tr>
<tr>
<td>Carpet fibre</td>
<td>0.60</td>
</tr>
<tr>
<td>High Tenacity Cotton type fibre</td>
<td>0.63–0.70</td>
</tr>
<tr>
<td>Technical yarn</td>
<td>0.72–0.90</td>
</tr>
<tr>
<td>Tyre-cord yarn</td>
<td>0.85–0.98</td>
</tr>
</tbody>
</table>
Polyester resins may contain 0.03 to 0.4 wt. % of titanium oxide (TiO\(_2\)) as delustering agent.

Table 1.8 shows the crystalline data of PET. Heat of fusion values reported for crystalline PET are generally in the range of 22.6 and 27.8 kJ/mol. Polyethylene glycol (PEG) with different molecular weights is added to produce small crystallites when crystallised from an amorphous solid, rather than forming one large single crystal. Light tends to scatter as it crosses the boundaries between crystallites and the crystalline PET is opaque and white in most cases. Fibre drawing is among the few industrial processes that produces a nearly single crystal product. PET fibre when exposed to elevated temperatures for 28 days in air, 90 and 30 per cent of strength retained at 150 and 200˚C respectively, whereas exposure for seven days in pressurised steam retains 90 per cent of strength at 100˚C and 0 per cent at 150˚C.

1.5.4 Applications

PET grades with relatively higher molecular weights are used for making industrial filaments, which can be either thick for rubber tyres, conveyor belts, seat belts, hoses and ropes, coated fabrics, etc. or relatively thinner for sewing threads, light-weight coated fabrics, etc. Staple fibre finds major use in making blended fabrics, low denier fibre for blending with cotton and coarser fibre for blending with wool. Fabrics made from PET POY

<table>
<thead>
<tr>
<th>a (nm)</th>
<th>b (nm)</th>
<th>c (nm)</th>
<th>α (°)</th>
<th>β (°)</th>
<th>γ (°)</th>
<th>(\rho_{\text{crystal}}) (g/cc)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.554</td>
<td>0.414</td>
<td>1.086</td>
<td>107.5</td>
<td>112.2</td>
<td>92.2</td>
<td>1.471</td>
<td>(Astbury, 1946)</td>
</tr>
<tr>
<td>0.456</td>
<td>0.594</td>
<td>1.075</td>
<td>98.5</td>
<td>118</td>
<td>112</td>
<td>1.455</td>
<td>(de Daubeny, 1954)</td>
</tr>
<tr>
<td>0.452</td>
<td>0.598</td>
<td>1.077</td>
<td>101</td>
<td>118</td>
<td>111</td>
<td>1.479</td>
<td>(Tomashpolskii, 1964)</td>
</tr>
<tr>
<td>0.448</td>
<td>0.585</td>
<td>1.075</td>
<td>99.5</td>
<td>118.4</td>
<td>111.3</td>
<td>1.515</td>
<td>(Fakirov, 1975)</td>
</tr>
<tr>
<td>0.444</td>
<td>0.591</td>
<td>1.067</td>
<td>100.1</td>
<td>117</td>
<td>111.8</td>
<td>1.515</td>
<td>(Asano, 1999)</td>
</tr>
<tr>
<td>0.452</td>
<td>0.592</td>
<td>1.070</td>
<td>99.8</td>
<td>117.5</td>
<td>111.4</td>
<td>1.479</td>
<td>(Hall, 1984a)</td>
</tr>
<tr>
<td>0.448</td>
<td>0.588</td>
<td>1.071</td>
<td>100.1</td>
<td>117.9</td>
<td>110.7</td>
<td>1.501</td>
<td>(Heuval, 1978)</td>
</tr>
<tr>
<td>0.450</td>
<td>0.590</td>
<td>1.076</td>
<td>100.3</td>
<td>118.6</td>
<td>110.8</td>
<td>1.501</td>
<td>(Kinoshita, 1979), (Kitano, 1995)</td>
</tr>
<tr>
<td>0.462</td>
<td>0.592</td>
<td>1.068</td>
<td>99.8</td>
<td>127.6</td>
<td>104.9</td>
<td>1.578</td>
<td>(Liu, 1997a)</td>
</tr>
</tbody>
</table>

Fibre grade is usually in the range of 0.40 to 0.98 dL/g and this fibre grade may contain 0.03 to 0.4 wt. % of titanium oxide (TiO\(_2\)) as delustering agent.
microfilaments are breathable and water-repellent with soft drape and pleasant feel.

1.6 Poly(trimethylene terephthalate) (PTT)

Poly(trimethylene terephthalate) (PTT) is otherwise known as poly(propylene terephthalate) (PPT). PPT nomenclature does not distinguish whether the glycol moiety is derived from branched 1,2-propane diol or linear 1,3-propane diol. The PTT abbreviation is more popular and widely used than PPT. It is also known as 3GT. The repeat unit molecular weight is 206. Futura Polyesters Ltd is the first company outside USA to produce PTT resin. Shell chemical company and DuPont’s PTT fibres are commercially known as Corterra™ and Sorona™ respectively. PTT is prepared from DMT (or) PTA and 1,3-propane diol (PDO). PTT has been a recent addition to the list of commercial aromatic polyesters, even though it was first synthesized by Whinfield and Dickson in 1941 (Whinfield and Dickson, 1946). PTT can be processed by extrusion, injection moulding, blow moulding and melt-spinning and the molten polymer has good melt strength. Chemical structure of PTT is as follows:

\[
\begin{array}{c}
\text{CH}_2 \text{CH}_2 \text{CH}_2 \text{O} \\
\text{O} \\
\text{C} \text{O} \text{CH}_2 \text{CH}_2 \text{CH}_2 \text{O}
\end{array}
\]

PTT had never gone beyond academic interest until recently, because PDO was very expensive initially, selling at about $20 per kg. PDO was first marketed by Degussa AG as a fine chemical and was synthesized from acrolein by selective hydration under pressure at 50°C using acid catalyst followed by hydrogenation using Raney Nickel as catalyst. Later, Shell invented an attractive less expensive route in which ethylene oxide is hydroformulated into 3-hydroxy propanol by using synthesis gas (a mixture of carbon monoxide and hydrogen) and cobalt as catalyst. Then 3-hydroxy propanol aqueous solution was concentrated and hydrogenated to produce PDO. Recently, DuPont has announced that PDO based on biochemical route using corn starch as feed stock will be commercialized in 2007.

1.6.1 Polymer production

Similar to PET production, PTT can be produced by polycondensation reaction through the DMT (or) PTA route. PTT can be produced by the
transesterification of PDO and DMT in the melt at 180–200°C using a zinc acetate dehydrate catalyst. After the completion of precondensation, the temperature was raised to 265°C and the melt was polymerised under vacuum of <0.3 mm of Hg using a titanium butoxide catalyst. Because of the lower reactivity of PDO, a more active catalyst such as titanium or tin is needed for PTT. Undesired side reactions occur, during both precondensation and polycondensation reactions, leading to the formation of acrolein, allyl alcohol, dipropyl glycol and cyclic dimer by-products. Among these acrolein is a very strong lachrymator, which can irritate lung and respiratory tracts and affect breathing. The US Occupational Safety and Health Agency industrial hygiene guidelines gave the time-weighted exposure limit of acrolein over a period of 8 h as 0.1 ppm, while the short-term exposure limit for 15 min is 0.3 ppm.

PTT should be dried to a moisture level of <30 ppm, in a close-loop hot air dryer having a dew point preferably lower than −40°C at 130°C for four hours. Otherwise, hydrolytic degradation will take place during melt processing. The dried polymer can be extruded at 250–270°C into bulk continuous filaments (BCFs), partially oriented yarn (POY), spin-draw yarn (SDY) and staple fibre. Because of its low Tg, PTT fibres and fabrics are disperse dyed at atmospheric boil without the need of a carrier (Traub et al., 1995). PTT dyed at atmospheric boil has very good colour fastness against light, ozone and NOx, similar to PET (Chuah et al., 1995; Yang et al., 1999).

1.6.2 Properties

PTT’s excellent stress recovery property is similar to that of nylon and this makes PTT an excellent candidate for textile applications. When compared with other polyesters, PTT has a very good tensile elastic recovery property, which descends in the following order: PTT > PBT > PET (Ward, 1976). PTT is not readily soluble in solvents, which are commonly used for amorphous PET, because of its rapidly crystallising nature. However, PTT is readily soluble in stronger solvents such as hexafluoroisopropanol (HFIPA) (or) in a 1:1 mixture of trifluoroacetic acid and dichloromethane at room temperature. With care, when heated to 110°C, PTT dissolves in a 60:40 mixture of phenol/terachloroethane (Chuah, 2001). The IV of fibre producing pellets is 0.92 dL/g and the corresponding molecular weight is 48700 and polydispersity is 2. PTT is extrusion spun. Like PET, PTT is sensitive to hydrolysis degradation and should be dried before processing. Unlike PET, there is no need to crystallise PTT chips before drying. A typical fibre grade PTT has a melt strength of about 200 Pa-sec at 260°C and at a shear rate of 200 per sec.
Contrary to the widely believed concept that aromatic polyesters with odd numbers of methylene units are more difficult to crystallise, PTT is fast crystallising. The rate of crystallisation is about an order of magnitude faster than PET. In terms of crystallising ability PTT is in between PET and PBT. Crystal density of PTT is 1.387 g/cc. Unit cell parameters are tabulated in Table 1.9. Unit cell of PTT is triclinic crystalline structure, each cell of which contains two chemical repeat units and the crystal space group is \( \rho I \) (Dandurand et al., 1979).

Hall has reviewed the crystal structures of PTT (Hall, 1984). Molecular configuration consists of rigid planar terephthaloyl moieties alternating with more flexible trimethylene moieties. The O – CH\(_2\) – CH\(_2\) – CH\(_2\) – O segment of the chain has a near trans-gauche-gauche-trans conformation. Studies by Ward (Ward et al., 1976) showed that PTT had an elastic unit cell, the (002) crystal lattice spacing of PTT responded immediately to the applied stress and the deformation was reversible below its critical strain. The microscopic reversible crystal chain deformation was attributed to the PTT’s three methylene units arranged in a highly contracted and a very compliant gauche-gauche conformation (Desborough et al., 1979, Poulin-Dandurand et al., 1979). PTT was found to have a very low X-ray crystal modulus of 2.59 GPa compared with 107 GPa of PET. The heat of fusion of 100 per cent crystalline PTT as measured by Pyda et al. (1998) is 30 kJ/mol, which is in good agreement with the values reported by Gonzalez et al. (1988a). Based on the DSC measurements, melting point is reported to be at 237°C (Pyda et al., 1998) and glass transition temperature is reported to be in the range of 37–45°C and melting point (Pyda et al., 1998; Gonzalez et al., 1988a) whereas higher glass transition temperature (59–69°C) was observed in dynamic mechanical analysis (DMA) studies. Other transitions observed are \( \beta \) at –70°C and \( \gamma \) at about –120°C (Gonzalez et al., 1988b).

### Table 1.9 Unit cell parameters of PTT

<table>
<thead>
<tr>
<th>Dimension (nm)</th>
<th>(Dandurand, 1979)</th>
<th>(Desborough, 1979)</th>
<th>(Yang, 2001)</th>
<th>(Tadokoro, 1979)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>0.4637</td>
<td>0.4620</td>
<td>0.453</td>
<td>0.458</td>
</tr>
<tr>
<td>b</td>
<td>0.6226</td>
<td>0.6200</td>
<td>0.6150</td>
<td>0.622</td>
</tr>
<tr>
<td>c</td>
<td>1.864</td>
<td>1.830</td>
<td>1.861</td>
<td>1.812</td>
</tr>
<tr>
<td>( \alpha ) (°)</td>
<td>98.4</td>
<td>98.0</td>
<td>96.87</td>
<td>96.9</td>
</tr>
<tr>
<td>( \beta ) (°)</td>
<td>93.0</td>
<td>90.0</td>
<td>92.21</td>
<td>89.4</td>
</tr>
<tr>
<td>( \gamma ) (°)</td>
<td>111.5</td>
<td>112.0</td>
<td>110.97</td>
<td>111.0</td>
</tr>
<tr>
<td>( \rho )(_{\text{crystal}} ) (g/cc)</td>
<td>1.387</td>
<td>1.427</td>
<td>1.448</td>
<td>1.43</td>
</tr>
</tbody>
</table>
1.6.3 Applications

PTT carpets (Chua, 1996) showed excellent resiliency in walk test experiments, equivalent to nylon and much better than both PET and polypropylene. The high resilience characteristic of PTT is due to its unique crystal structure, which is very spring-like in its long axis. Further, it had a lower static charge of $<3.5 \text{kV}$, and was resistant to coffee, mustard, butadiene, red acid dyes and other stains (Chua, 1996). PTT is used in ready-to-wear, stretch apparels (Heschmeyer, 2000), active-wear, intimate apparels and inner linings, automotive and home upholstery. PTT monofilaments can be used to make paper forming fabrics, which find use in papermaking machines (Hsu, 1992). PTT is also used to make non-woven (Asahi Chemical Industry Co. Ltd, 1999) fibres, artificial leather (Goto and Muraoka, 1999), zip fasteners (Kawase and Kuratsuji, 1976), umbrella fabric (Yamamoto, 1999), racket guts and musical bowstrings (Oue and Yamaziki, 2000), pantyhose (Hiraga and Sonoda, 1999), cheese packaging (Kato and Fujimoto, 1999), hook-and-loop fasteners (Ohira, 1999), magnetic recording discs (Hosoi, 1988), electrical connectors (Hironaka and Suzuki, 1999) and flexible transparent film.

1.7 Poly(butylene terephthalate) (PBT)

Poly(butylene terephthalate) (PBT) belongs to aromatic polyester family. It is otherwise known as polytetramethylene terephthalate or 4GT. This thermoplastic polyester is prepared by polycondensation of 1,4-butane diol (BDO) with PTA (or) DMT. The repeat unit molecular weight is 220. PBT was brought to market by Celanese in late 1960s. Structure of PBT is given below:

\[
\begin{array}{c}
\text{O} \\
\text{C--O---CH}_2\text{---CH}_2\text{---CH}_2\text{---CH}_2\text{---O} \\
\text{n}
\end{array}
\]

PBT differs only slightly from PET in chemical composition but more substantially in its properties such as higher crystallisation rate and lower melting point. PBT fibres are manufactured by Zimmer and Ticona in the name of Celanex®. PBT resin is similar to polyamide resin nylon 6 in many aspects such as: (i) ability to crystallise quickly; (ii) good solvent resistance and mechanical properties; and (iii) similar melting point. PBT is different from nylon 6 with respect to moisture absorption. PBT absorbs less moisture than nylon. PBT shows more resistance to staining than nylon and can...
be coloured by the use of pigments. However PBT is more difficult to
colour by solution dying than nylon.

1.7.1 Polymer production

PBT production is similar to PET and can be produced by polycondensation reaction from DMT (or) PTA and BDO. In PBT production, THF is formed as by-product by irreversible acid-catalysed dehydration of BDO. Alternatively, THF is produced by back-biting process. DMT route produces approx. 6 mol. % and PTA route 13 mol. % of THF (Schumann, 1990). Formation of THF is minimised by the addition of 2–8 per cent of water to PTA and BDO (US Patent 3,936,421), the usage of combination of titanium and tin catalysts (US Patent 4,014,858), the usage of titanium catalyst (US Patent 5,015,759), the slow addition of PTA (US Patent 4,329,444), or the addition of a portion of BDO at a later stage of reaction (US Patent 4,565,241) or carrying out polycondensation before PTA is exhausted (US Patent 4,364,213). The other by-product formed is 1,3-butadiene. In melt phase, $M_n$ of 20000 to 35000 is achieved and is used for both fibre and engineering plastics applications. SSP is used to increase the molecular weight to 40000, which is suitable for certain injection moulding and extrusion applications. After polycondensation, i.e., before SSP, PBT resin is already crystalline and opaque and therefore, unlike PET, there is no need to precrystallise PBT resin in a fluid bed. Due to lower melting point (225°C), SSP is carried out at 180 to 200°C. Even though PBT is less sensitive to oxidation than PET, SSP is carried out in nitrogen. Reaction rate of PBT is faster than that of PET, IV is increased from 0.8 to 1.2 dL/g in about 12 hours.

1.7.2 Properties

PBT displays good solvent resistance, high heat resistance, good elongation, high strength and modulus, excellent electrical properties, high gloss, good inherent lubricity and wear resistance. Density of amorphous PBT is in the range of 1.265–1.268 g/cc and that of crystalline is 1.395 g/cc. Melting point and glass transition temperature of PBT are 225°C and 25°C respectively. Heat of fusion for crystalline PBT is 32 kJ/mol (Cheng et al., 1988). Under moderate conditions, the degree of crystallinity is about 35 to 40 per cent. Water absorption is less than 0.1 per cent after 24 h. However, PBT is not recommended for extended use in water/aqueous solution above 52°C. PBT is intrinsically resistant to detergents, weak acids and bases, aliphatic hydrocarbons, fluorinated hydrocarbons, alcohols, ketones, MEG, carbon tetrachloride, oils and fats at ambient temperature.
Among the aromatic polyesters, PBT has high crystallisation rate next to PBN, about an order of magnitude faster than PTT, which in turn is an order of magnitude faster than that of PET (Chuah, 2001). Similar to PTT, PBT has triclinic crystalline structure. Two types of triclinic structures exist namely α- and β-form, which are reversible. Unit cell parameters of α-form and β-form of PBT are set out in Tables 1.10 and 1.11. In the non-extended state, the α-form prevails with gauche-trans-gauche conformation of butylenes moiety. The β-form exists in the extended state with all trans conformation depending on the drawing condition. Unit cell volume of α- and β-forms of PBT are 0.261 and 0.267 nm³.

### Table 1.10 Unit cell parameters of α-form of PBT

<table>
<thead>
<tr>
<th>PBT Dimension</th>
<th>α-form (Bornschlegl, 1980)</th>
<th>α-form (Hall, 1976)</th>
<th>α-form (Joly, 1975)</th>
<th>α-form (Liu, 1997)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a (nm)</td>
<td>0.482</td>
<td>0.489</td>
<td>0.487</td>
<td>0.494</td>
</tr>
<tr>
<td>b (nm)</td>
<td>0.593</td>
<td>0.595</td>
<td>0.596</td>
<td>0.598</td>
</tr>
<tr>
<td>c (nm)</td>
<td>1.174</td>
<td>1.167</td>
<td>1.171</td>
<td>1.156</td>
</tr>
<tr>
<td>α (°)</td>
<td>100</td>
<td>98.9</td>
<td>100.1</td>
<td>99.8</td>
</tr>
<tr>
<td>β (°)</td>
<td>115.5</td>
<td>116.6</td>
<td>116.6</td>
<td>116.5</td>
</tr>
<tr>
<td>γ (°)</td>
<td>111</td>
<td>110.9</td>
<td>110.3</td>
<td>111.15</td>
</tr>
<tr>
<td>ρ&lt;sub&gt;crystal&lt;/sub&gt; (g/cc)</td>
<td>1.403</td>
<td>1.392</td>
<td>1.396</td>
<td>1.397</td>
</tr>
</tbody>
</table>

### Table 1.11 Unit cell parameters of β-form of PBT

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>a (nm)</td>
<td>0.473</td>
<td>0.473</td>
<td>0.473</td>
<td>0.469</td>
</tr>
<tr>
<td>b (nm)</td>
<td>0.575</td>
<td>0.583</td>
<td>0.588</td>
<td>0.580</td>
</tr>
<tr>
<td>c (nm)</td>
<td>1.311</td>
<td>1.290</td>
<td>1.306</td>
<td>1.300</td>
</tr>
<tr>
<td>α (°)</td>
<td>104.2</td>
<td>101.9</td>
<td>103.3</td>
<td>101.9</td>
</tr>
<tr>
<td>β (°)</td>
<td>120.8</td>
<td>119.4</td>
<td>119.8</td>
<td>120.5</td>
</tr>
<tr>
<td>γ (°)</td>
<td>100.9</td>
<td>105.1</td>
<td>104.4</td>
<td>105</td>
</tr>
<tr>
<td>ρ&lt;sub&gt;crystal&lt;/sub&gt; (g/cc)</td>
<td>1.33</td>
<td>1.33</td>
<td>1.32</td>
<td>1.36</td>
</tr>
</tbody>
</table>

1.7.3 Applications

PBT is used for textile applications due to its stretchability and improved dyeability. PBT fibres are used as tooth brush bristles, carpet yarn and sports wear, and in apparel, underwear and hosiery. It is also used for swim
wear since it has high tenacity, stability and good resistance to chlorine. PBT is preferably used for the production of engineering plastics due to its combination of dimensional stability, tensile strength, increased flexibility and fast crystallisation rate. PBT resins are used in drapery hardware, pen barrels, heavy duty zippers, hair dryers, pocket calculators, iron and toaster housings and food processor blades.

1.8 Poly(1,4-cyclohexylene dimethylene terephthalate) (PCT)

Poly(1,4-cyclohexylene dimethylene terephthalate) is abbreviated as PCT. It is prepared from 1,4-cyclohexanediethanol (CHDM) and PTA (or) DMT. CHDM imparts the following advantages for polyester resins: (i) higher Tg; (ii) higher reactivity; (iii) glossiness and transparency; and disadvantages for polyester resins such as: (i) higher viscosity; (ii) lower solvent solubility; and (iii) lower tendency to crystallise. Chemical structure of PCT is as shown below:

Unlike other glycols, CHDM exists in cis and trans forms. The cis form imparts increased gas barrier property when compared to trans form. PCT is less dense than PET so it offers great bulk. Specific gravity of PCT is 1.23 whereas for PET it is 1.38. PCT provides softness inherent to the molecular composition. PCT offers better release properties than PET which means that it is reusable and washes and cleans up easily. PCT has better hydrolytic stability than PET so it withstands better the rigours of repeated washing and sterilization. PCT has a higher melting point than PET (290 vs. 250°C). PCT has higher glass transition temperature than PET (88 vs. 80°C). Heat distortion temperature of PCT is 75 and 65°C at 0.455 and 1.82 MPa respectively. PCT has better chemical resistance to auto fluids and printed circuit board cleaning chemicals than PET. PCT is sold under the trade name of Kodel II by Eastman Chemical Products Inc. PCT fibres can be used for high temperature applications, for example autoclave and industrial applications such as filtering, insulation, etc. Because of its superior softness with resilience, PCT fibres can be used for pillows, cushions, bed pads, carpets, etc. The safe ironing temperature for
PCT based clothes is 205°C whereas for PET it is 145°C. PCT fibres shrink less than PET fibres. Percentage shrinkage at 190°C (air) are <1 and <6 for PCT and PET fibres respectively. Percentage shrinkage at 100°C (water) for PCT fibres is <0.5 and for PET fibres is <1. PCT fibre has better resiliency than PET, without compromising high temperature resistance. It competes successfully against nylon in the carpet yarn market. The higher HDT has allowed PCT to carve out a niche market in surface mount components, electrical connectors, sensors and switches, especially those which require IR oven compatibility.

1.9 Poly(ethylene 2,6-naphthalate) (PEN)

The expansion of PEN is poly(ethylene 2,6-naphthalate) and is three times more expensive than its analogue, PET. The systematic nomenclature of PEN is poly(oxy-1,2-ethanediylxocarbonyl-2,6-naphthalenedicarbonyl). Several other poly(ethylene naphthalate)s such as poly(ethylene naphthalene 1,8-dicarboxylate) (Sakai et al., 1992; Yokota, 1994), poly(ethylene naphthalene 2,7-dicarboxylate) (Khanna et al., 1994; Sullivan and Hoyt, 1996), poly(ethylene naphthalene 1,5-dicarboxylate) (White et al., 1999) are reported. NDC has been commercially available from British Petroleum since 1996. Teijin Ltd of Japan has manufactured PEN for its own uses in films since 1973. In the late 1980s, both Eastman Chemicals and Goodyear developed PEN. Eastman’s PEN was called 5XO (five times better oxygen barrier than PET) and Goodyear’s PEN was called HP (high performance). Performance Fibers manufactures PEN fibres under the name of PenTec®. Futura Polyesters Ltd makes PEN with different IVs ranging from 0.45 to 0.85 dL/g. The chemical structure of PEN is:

\[
\begin{align*}
\text{CH}_2\text{O} & \quad \text{O} \\
\text{C} & \quad \text{O} \\
\text{C} & \quad \text{O} \\
\text{CH}_2 & \quad \text{CH}_2 & \quad \text{O} \\
\end{align*}
\]

NDA/NDC can be synthesized from various raw materials (Shikkena and Wheaton, 1989, 1990; Jia and Sheng, 1999; Motoyuki et al., 1999; Holzhauer and Young, 1993; Uchida and Marumo, 1989) as follows:
(a) from \( \alpha \)-xylene and 1,3-butadiene

\[
\begin{align*}
\text{CH}_3 & \quad \text{CH}_3 \\
\text{CH}_3 & \quad \text{CH}_2=\text{CH}-\text{CH}=\text{CH}_2 \\
\text{CH}_3 & \quad \text{CH}_2=\text{CH}-\text{CH}_2\text{CH}_3 \\
\text{H}_2 &
\end{align*}
\]

(b) from 2-methylnaphthalene and acetyl acetonate

\[
\begin{align*}
\text{CH}_3 & \quad \text{O} \quad \text{O} \\
\text{CH}_3 & \quad \text{CH}_3-\text{C}-\text{O}-\text{C}-\text{CH}_3 \\
\text{BF}_3, \text{HF} & \quad \text{CH}_3-\text{C}-\text{O} \\
\text{Co, Mn} & \quad \text{Br/O}_2 \\
\text{H}_3\text{CO}-\text{C} & \quad \text{O} \\
\text{CH}_3\text{OH} & \quad \text{C}-\text{OH} \\
\text{NDA} & \quad \text{NDC}
\end{align*}
\]
(c) from naphthalene and propylene

\[
\begin{align*}
\text{NDC} & \quad + \quad \text{CH}_3 - \text{CH} = \text{CH}_2 \\
& \quad \xrightarrow{\text{Catalyst}} \\
\text{NDA} & \quad \xrightarrow{\text{O}_2, \text{Catalyst}} \\
\end{align*}
\]

(d) from potassium naphthalene carbonate

\[
\begin{align*}
\text{COOK} & \quad \xrightarrow{\text{CO}_2} \\
\text{HOOC} & \quad \xrightarrow{\text{HCl}} \\
\text{CH}_3\text{OOC} & \quad \xrightarrow{\text{Catalyst}} \\
\end{align*}
\]

(e) from 1,8-naphthalene dicarbonate

\[
\begin{align*}
\text{COOK} & \quad \xrightarrow{\text{KOOC}} \\
\text{COOH} & \quad \xrightarrow{\text{HCl}} \\
\text{CH}_3\text{OOC} & \quad \xrightarrow{\text{Catalyst}} \\
\end{align*}
\]
1.9.1 Polymer production

PEN can be prepared from MEG and NDC (or) NDA by polycondensation reaction. Alternatively, PEN can be prepared by ROP (Hubbard and Brittain, 1996) of cyclic PEN in the presence of a tin catalyst. Cyclic PEN is prepared by the reaction of di(chlorocarbonyl)-2,6-naphthalene with MEG (Brunelle and Bradt, 1993; Brunelle, 1994) in dichloromethane in the presence of diazabicyclo[2.2.2]octane.

1.9.2 Properties

Both glass transition and melting temperatures of PEN are higher than that of PET. PEN also exhibits an increased tensile strength, as well as elastic modulus and elongation at break. Both oxygen and carbon dioxide gas barrier properties are about 70–80 per cent increased compared with PET. Due to high melt viscosity, PEN needs a higher processing temperature during the melt extrusion process and is slow crystallising resin. Density of PEN is lower than PET.

Depending on the previous thermal history, PEN has two crystalline structures namely α- (Mencik, 1967) and β-forms (Buchner et al., 1989), which are triclinic. When crystallisation temperature is ≤200°C, α-crystal is formed whereas the formation of β-crystal is at crystallisation temperature of ≥240°C. Their densities are 1.407 and 1.439 g/cc and that of amorphous material is 1.326 g/cc (Liu et al., 1998). The α-form crystal structure has dimensions of \( a = 0.651 \) nm, \( b = 0.575 \) nm, \( c = 1.32 \) nm, \( α = 81.33^\circ \), \( β = 144^\circ \) and \( γ = 100^\circ \) whereas β-form crystal structure has dimensions of \( a = 0.926 \) nm, \( b = 1.559 \) nm, \( c = 1.217 \) nm, \( α = 121.6^\circ \), \( β = 95.57^\circ \) and \( γ = 122.52^\circ \).

The dielectric constant (\( ε \)) of PEN is lower than that of PET in the entire temperature range. PEN is more suitable for high temperature applications because tanδ of PEN is higher than that of PET at room temperature, but is lower than those of PET at 90 and 130°C. PEN has greater chemical resistance than PET. Even though this is a desirable property for many applications, it makes PEN difficult to characterise by IV measurements, GPC, NMR and end-group analysis and this difficulty is even greater for highly crystalline material. Due to the extended chromophore group naphthalene, PEN absorbs UV light up to 380 nm whereas PET’s absorption is up to 313 nm. This property helps to use PEN in packaging applications. A blue-white visible fluorescence results when PEN is excited by black light emitting at ~360 nm. This permits ready identification and separation of PEN from PET chips, containers, films and fibres (Ouchi et al., 1976).

PEN begins to stick at 140°C as the Tg is at 121°C. But the crystallisation rate is not significant below 180°C and the maximum rate of crystallisation...
is at 200°C. However, at a crystallisation temperature of about 200°C, it has a strong tendency to stick strongly in the presence of high moisture content. PEN expands and bursts; known as ‘popcorning’. PEN requires 22 h at 235°C, to increase IV from 0.55 to 0.75 dL/g. The Mark–Houwink relationship for PEN resin is as follows:

\[
[\eta] = 2.317 \times 10^{-3} \times M_w^{0.514}
\]

In the above relationship, the IV solvent was used and 60:40 phenol/tetrachloroethane at 30°C.

1.9.3 Applications

PEN monofilaments are used for paper making felts, where the higher modulus, temperature and hydrolysis resistance are required. PEN fibre is used to make high performance sail cloth in racing, in tyre cord and also for making narrow and broad woven fabrics. PEN has been successfully used to contain liquid anaesthetics.

1.10 Polylactic acid (PLA)

Polylactic acid (PLA) differs from the previously discussed polyesters because PLA is based on renewable resources and is a biodegradable polymer. It is otherwise known as polylactide. The chemical structure of PLA is shown below:

\[
\begin{array}{c}
\text{O} \\
\text{C} \\
\text{CH} \\
\text{CH}_3
\end{array}
\]

It is produced from any starch or sugar containing raw material such as corn, cereals, cassava, sugar cane, etc. PLA can be used as packaging materials such as biaxially oriented films, extrusion sheets for thermoforming and plastic bottles, textile materials such as filaments, staple fibres and non-wovens and engineering plastics. Although PLA is expensive it will become cheaper than PET, now that the crude oil price has risen above US$80 per barrel. Depending upon the content of L and D isomers in PLA, melting point ranges from about 150°C to 210°C. Even though the polymer chain contains no aromatic rings, the steric hindrance of the pendant methyl groups gives some rigidity and thermal stability. Fibre elasticity and resiliency of PLA is similar to PTT or PBT and is attributed to the partial coiling of the backbone chain in the crystalline form.
1.10.1 Production of PLA

Production consists of two stages. During the first stage, glucose is fermented to produce lactic acid and in the second stage, lactic acid is polymerized to PLA. Fermentation of glucose produces not only lactic acid, in the form of sodium lactide but also numerous impurities such as proteins, cellular mass, etc. Sodium lactide is purified and converted into lactic acid. For polymerizing lactic acid, excellent purity is a requisite. Lactic acid undergoes polycondensation to form oligomer, which is thermally depolymerised to cyclic dimer namely dilactide. ROP of dilactide produces PLA and about 5 per cent of non-convertible dilactide, which is removed by applying a vacuum. Otherwise, rapid hydrolytic degradation takes place.

1.11 World market

Many manufacturers of raw materials have already installed (or are planning to schedule) their manufacturing units and the list is given in Tables 1.12 and 1.13 for PTA and MEG respectively. Many polyester manufacturing plants have started production recently and many more are scheduled to start production in the future. These manufacturers are listed in Table 1.14. Global polyester capacity greatly exceeds consumption. With the

<table>
<thead>
<tr>
<th>Company</th>
<th>Location</th>
<th>Technology</th>
<th>Capacity KTA</th>
<th>Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indian Oil</td>
<td>India</td>
<td>Invista</td>
<td>600</td>
<td>2006</td>
</tr>
<tr>
<td>Mitsubishi Chemical</td>
<td>China/India</td>
<td>MCC</td>
<td>600/800</td>
<td>2007/2009</td>
</tr>
<tr>
<td>Yisheng</td>
<td>China</td>
<td>Invista</td>
<td>600</td>
<td>2005</td>
</tr>
<tr>
<td>SMPC</td>
<td>Thailand</td>
<td>Mitsui</td>
<td>500</td>
<td>2005</td>
</tr>
<tr>
<td>Indorama</td>
<td>Thailand</td>
<td>Invista</td>
<td>640</td>
<td>2006</td>
</tr>
<tr>
<td>NPC</td>
<td>Iran</td>
<td>MCC</td>
<td>350</td>
<td>2006</td>
</tr>
<tr>
<td>OPC (FET)</td>
<td>China</td>
<td>Invista</td>
<td>600</td>
<td>2006</td>
</tr>
<tr>
<td>Reliance</td>
<td>India</td>
<td>Invista</td>
<td>600</td>
<td>2006</td>
</tr>
<tr>
<td>Yangzi</td>
<td>China</td>
<td>Invista</td>
<td>600</td>
<td>2007</td>
</tr>
<tr>
<td>BP Zhuhai</td>
<td>China</td>
<td>BP</td>
<td>900</td>
<td>2008</td>
</tr>
<tr>
<td>DaHua</td>
<td>China</td>
<td>Hitachi/Tuntex</td>
<td>1500</td>
<td>2010</td>
</tr>
<tr>
<td>Zinan</td>
<td>China</td>
<td></td>
<td>530</td>
<td>2009</td>
</tr>
<tr>
<td>Liaoyang</td>
<td>China</td>
<td>Invista</td>
<td>600</td>
<td>2006</td>
</tr>
<tr>
<td>Taekwang</td>
<td>Korea</td>
<td>Technimont</td>
<td>500</td>
<td>2007</td>
</tr>
<tr>
<td>Xianglu</td>
<td>China</td>
<td>Hitachi/Tuntex</td>
<td>1500</td>
<td>2009</td>
</tr>
<tr>
<td>Jialong</td>
<td>China</td>
<td>Invista</td>
<td>600</td>
<td>2009</td>
</tr>
<tr>
<td>Hanbang</td>
<td>China</td>
<td>Invista</td>
<td>600</td>
<td>2010</td>
</tr>
</tbody>
</table>
In 1997, the crude oil price was about $25/bbl and since then the price has passed $80/bbl. In the late 1990s, the polyester price was about $800/MT and had risen to $1200/MT at the time of writing. The tripling of the crude oil price has not proportionally increased the polyester price and the increase is only 50 per cent. The newly constructed fully integrated plants and increased competition among the polyester manufacturers has reduced the overheads and squeezed the profit of polyester manufacturers.

Many polyester industries are expected to emerge in the next couple of years, from the Middle East, China and India. Further polyester industry

---

### Table 1.13 Some of the new manufacturers of MEG in Asia

<table>
<thead>
<tr>
<th>Company</th>
<th>Location</th>
<th>Capacity KTA</th>
<th>Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>ShangHaiPC</td>
<td>China</td>
<td>320</td>
<td>2007</td>
</tr>
<tr>
<td>Nanya</td>
<td>Taiwan</td>
<td>700</td>
<td>2007</td>
</tr>
<tr>
<td>NPC</td>
<td>Iran</td>
<td>800</td>
<td>2007</td>
</tr>
<tr>
<td>TOC</td>
<td>Thailand</td>
<td>100</td>
<td>2008</td>
</tr>
<tr>
<td>LiaoYang</td>
<td>China</td>
<td>200</td>
<td>2008</td>
</tr>
<tr>
<td>Aramco</td>
<td>Saudi</td>
<td>600</td>
<td>2008</td>
</tr>
<tr>
<td>Sharq</td>
<td>China</td>
<td>700</td>
<td>2008</td>
</tr>
<tr>
<td>Lotte Daesan</td>
<td>Korea</td>
<td>400</td>
<td>2008</td>
</tr>
<tr>
<td>Equate</td>
<td>Kuwait</td>
<td>600</td>
<td>2009</td>
</tr>
<tr>
<td>Yansab</td>
<td>Saudi</td>
<td>700</td>
<td>2009</td>
</tr>
<tr>
<td>Shell</td>
<td>Singapore</td>
<td>750</td>
<td>2009</td>
</tr>
</tbody>
</table>

### Table 1.14 Some of the recent PET manufacturers and their capacity

<table>
<thead>
<tr>
<th>Company</th>
<th>Location</th>
<th>Capacity KTA</th>
<th>Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCH (Neo Gr)</td>
<td>Lithuania</td>
<td>150</td>
<td>2005</td>
</tr>
<tr>
<td>FET</td>
<td>China</td>
<td>160</td>
<td>2006</td>
</tr>
<tr>
<td>Wellman</td>
<td>USA</td>
<td>130</td>
<td>2006</td>
</tr>
<tr>
<td>Senezh (Europlast)</td>
<td>Russia</td>
<td>80</td>
<td>2006</td>
</tr>
<tr>
<td>Invista</td>
<td>Mexico</td>
<td>210</td>
<td>2006</td>
</tr>
<tr>
<td>Indorama</td>
<td>Lithuania &amp; USA</td>
<td>175 &amp; 80</td>
<td>2006</td>
</tr>
<tr>
<td>DAK</td>
<td>USA</td>
<td>200</td>
<td>2007</td>
</tr>
<tr>
<td>Voridian</td>
<td>USA</td>
<td>350</td>
<td>2007</td>
</tr>
<tr>
<td>NPC</td>
<td>Iran</td>
<td>140</td>
<td>2007</td>
</tr>
<tr>
<td>M &amp; G</td>
<td>Brazil</td>
<td>450</td>
<td>2007</td>
</tr>
<tr>
<td>Sabic</td>
<td>Saudi Arabia</td>
<td>250</td>
<td>2007</td>
</tr>
<tr>
<td>Sanfangxang</td>
<td>China</td>
<td>300</td>
<td>2007</td>
</tr>
<tr>
<td>W. B. Radici</td>
<td>China</td>
<td>200</td>
<td>2008</td>
</tr>
</tbody>
</table>
growth is limited by low returns; and escalating construction costs and salaries due to a shortage of experienced and skilled manpower. Even with the above constraints, polyester capacity is expected to grow from 18.8 million MT in 2007 to 22.7 million MT in 2010. About 4 to 5 per cent growth is estimated during the next five years in the textile industry and about 8 per cent in packaging applications.

1.12 Future trends

Among polyesters, PET will continue to be in the leading position for applications in packaging and textiles with more applications in ‘Technical Textiles’ in sectors of agriculture, building, geo, home, medical, packaging, etc. Also there will be more developments in textile applications for comfort, sportswear, etc. The global production of technical fibres is growing by 13 per cent with applications in the aerospace industry including space and defence. Quality improvements such as increased barrier capabilities, softness and loft will be further improved. An increased demand for renewable raw materials and disposable materials for environmental and cost reasons will be a central issue in the future. Growth in today’s booming markets will be influenced by a shortage of raw materials and energy. Polyester will continue to find newer applications because of its undisputable performance and properties.

At the current consumption rates, the current world crude oil reserves will be exhausted by the year 2043, unless new oil stocks are discovered. If, as predicted, the global population doubles in the next 50 years, the requirement for polyester resin at the current usage rate will also double. This will have a huge bearing on crude oil consumption. Therefore, the future of the polyester industry, in particular commodity resin, depends on how effectively the industry recycles polyester scrap and how quickly the industry moves from oil-based resources to renewable resources. The effective recycling of polyester resin will also reduce carbon dioxide emission, which in turn will minimise global warming. Even though, with the available technology today, one can make PDO and PLA using renewable resources, commercial production and effective utilisation have not yet taken full effect. PLA will become cheaper only when the crude oil price goes above $80/bbl. Till then, PLA will be only of academic interest, unless new technology is invented to make polyesters from renewable resources.

1.13 Acknowledgements

We acknowledge Mr S. B. Ghia, Chairman and Managing Director; Mr M. D. Dalal, Joint Managing Director and Mr S. Rangarajan, COO and
Executive Director of Futura Polyesters Ltd for giving permission to write this chapter.

1.14 Sources of further information and advice


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2

Polyamide fibers

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New Delhi, India

2.1 Introduction

Nylon is a generic word representing a class of polymers as polyamides. Nylon was a product of a basic research started by Wallace Corothers at DuPont in 1928. The polymer age really started when the new synthetic nylon fiber was introduced in the market in 1938. Nylon captured public imagination, as four million pairs of nylon stockings were sold in the first few hours of sale on May 15, 1940. During World War II, nylon fabrics were used as waterproof tents, lightweight parachutes among other things. In the process an important principle was established that it is possible to chemically link hundreds of simple molecules of a kind or more, to make polymers with unique set of properties. One interesting example is the ability of molten polymer undergoing large deformations, which allows continuous production of fine filaments from thicker extrudates and this property of deformation is in turn connected to cumulative interactions of long chain polymeric molecules.

The word nylon was introduced to signify the fineness of the filament i.e. a pound of nylon could be converted to the length equal to the distance between New York (NY) and London (LON). A variety of polyamides are presently manufactured and marketed under several different trade names. Among them nylon 6 and nylon 66 are the two most manufactured polyamides. There are several other nylons such as nylon 46, nylon 610, nylon 612, nylon 10, and nylon 12 which are used for specialized applications.

In general, there are two types of nylons, i.e. nylon XY, and nylon Z. Nylon 66 and nylon 6 are the examples of these two types respectively. In nylon XY type, the X refers to the number of carbon atoms in the diamine monomer, whereas, Y represents the number of carbon atoms in diacid monomer. In nylon Z type, Z refers to the number of carbon atoms in the monomer.
2.2 Nylon 66

Nylon 66 typenylons are synthesized from a diacid and diamine. For nylon 66, hexamethylene diamine (H) and adipic acid (A) are reacted to form H-A salt. The salt is polymerized to produce nylon 66 in the following manner.

\[
\begin{align*}
\text{NH}_2(CH_2)_6NH_2 + \text{COOH}(CH_2)_4\text{COOH} & \rightarrow \\
H[-\text{NH}(CH_2)_6\text{NHCO}(CH_2)_4\text{CO}]\text{OH} + \text{H}_2\text{O} \\
n H[-\text{NH}(CH_2)_6\text{NHCO}(CH_2)_4\text{CO}]\text{OH} & \rightarrow \\
H[-\text{NH}(CH_2)_6\text{NHCO}(CH_2)_4\text{CO}–]_n\text{OH} + (n - 1)\text{H}_2\text{O}
\end{align*}
\]

The purity of raw material is important in the polymer production and the specifications of these raw materials have been reported [1]. Hexamethylene diamine should be in the crystalline form with melting point of 40°C and boiling point of 204°–205°C. Adipic acid is in the white crystalline form with acid number of 368, with melting and boiling point of 152° and 216°C. The color, iron and water content are also controlled.

In the production process, the two monomers of nylon 66 are taken in the molar ratio of 1 : 1. These two components are reacted in methanol at elevated temperature, to obtain the salt which precipitates from methanol solution. The H-A salt is dissolved in water at concentration of 60%. The solution is heated to around 250°C for polycondensation for production of nylon 66. Loss of water in the process of polymerization should be taken into account for the net recovery of polymer [2]. Nylon 66 was earlier produced via a batch process with intermediate H-A salt. However, now the continuous process of production is favored for conversion of monomer to polymer.

To achieve desired molecular weights of the polymers, stabilization processes are used. This is achieved by taking slightly different than equivalent quantities of diacids and diamines. Say an excess of 1 percent of diacids is taken during the polymerization process, at a point of time, all long chain molecules will have carboxylic acid groups at both ends and polymerization does not proceed further. Thus an excess of one of the monomers decides the range of molecular weights achieved in the polymer. An equimolar ratio of 1 diamine to 1.015 diacid gives a polymer with a number average molecular weight of ~18000. This can also be achieved by adding say 1% of mono-functional reagent, such as acetic acid to equimolar quantities of diacids and diamines. Another method to make polyamides could be by using amino acids with one end as an amino group and another end as carboxylic acid. Typically, diamine and diacids are preferred for production of nylons as it is easier and cheaper to make these monomers compared to amino acids. There is no difficulty in making the monomers for making
the typical polyamides and usually there are several different routes to select one, depending on the price and availability of raw materials.

2.2.1 Monomer synthesis for nylon 66

Adipic acid

The principal commercial processes are based on the oxidation of cyclohexane, which usually proceeds in two stages. The first step entails oxidation with air, yielding either a mixture of cyclohexanone and cyclohexanol or predominantly cyclohexanol. These reaction products are oxidized in second stage with nitric acid to adipic acid. This process employs a soluble cobalt oxidation catalyst, reaction temperatures in the range of 150°–160°C, pressures between 800 to 1000 kPa, and catalyst concentrations of 0.3–3 ppm.

An interesting approach involves a two-step carbonylation of butadiene. In the first step butadiene is reacted with carbon monoxide and methanol in the presence of dicobaltoctacarbonyl and heterocyclic structure containing a tertiary nitrogen moiety (pyridines, picolines, quinolines, isoquinoline):

\[
\text{CH}_2=\text{CHCH}═\text{CH}_2 + \text{CO} + \text{CH}_3\text{OH} \rightarrow \\
\text{CH}_3\text{CH}═\text{CHCH}_2\text{COOCH}_3
\]

This reaction is carried out at 120°C and at pressure of 60 MPa. After removal of unreacted butadiene, the second step carbonylation is carried out at about 185°C and 3 MPa:

\[
\text{CH}_3\text{CH}═\text{CHCH}_2\text{COOCH}_3 + \text{CO} + \text{CH}_3\text{OH} \rightarrow \\
\text{H}_3\text{COOC(CH}_2)\text{4COOCH}_3
\]

The dimethyl adipate is isolated by distillation and converted to adipic acid by hydrolysis.

Hexamethylene diamine

The commercial processes for manufacture of hexamethylene diamine entail hydrogenation of adiponitrile. It is a continuous liquid phase process that is usually conducted at 75°C and 3 MPa pressure in the presence of chromium containing Raney nickel catalyst and aqueous sodium hydroxide:

\[
\text{NC(CH}_2)\text{4CN} \rightarrow \text{H}_2\text{(3Mpa), 75°C, Cr/Ni, (NaOH, H}_2\text{O)} \rightarrow \text{H}_2\text{N(CH}_2)\text{6NH}_2
\]
A newer DuPont process is characterized by the direct addition of HCN to butadiene. It is essentially a two stage process. The first step is a vapor phase operation. A gaseous mixture of butadiene, hydrogen cyanide, nitrogen, and hydrochloride in a ratio of 1:1:1:0.1 is contacted at about 215°C and atmospheric pressure with a copper magnesium chromite fixed bed catalyst for about 50 min:

$$CH_2=CHCH=CH_2 + HCN \rightarrow CH_3CH=CHCH_2CN + \Delta 2 \text{ and } \Delta 4 \text{ isomers}$$

The second step entails catalytic isomerization of the 2- and 3-pentene nitriles to 4-pentene nitrile, and the addition of a second mole of HCN:

$$2\text{- and } 3\text{-pentene nitriles } \rightarrow CH_3CH=CHCH_2CN + HCN \rightarrow NC(CH_2)_3CN$$

### 2.3 Nylon 6

Nylon 6 is a nylon Z type of polymer, where the Z represents the number of carbon atoms in the monomer. Nylon 6 is typically produced from caprolactam in the following manner.

$$n \left[HN-\left(CH_2\right)_5-CO\right] \rightarrow H\left[-HN-\left(CH_2\right)_5-CO-\right]_n OH$$

The purity of the caprolactam is critically assessed for the production of nylon 6 polymer. The specificities of the monomer are freezing point 69°C; permanganate number 71; molar ratio %; iron content as 0.5 ppm. Detailed analytical methods for purity analysis are reported in literature [3–5].

Caprolactam requires a catalyst, which converts small amounts of caprolactam to ε-aminocaproic acid which helps in polymerization process. The catalyst could be water, where the control of reaction is easy. Acid or base catalyst system may also be used, where the base catalyst system has the advantage of high production rate, high molecular weight, although the control of reaction in the system is rather critical. Some characteristics of catalyst system [6] are given in Table 2.1.

#### 2.3.1 Molecular weight in water catalyst system

In a water catalyst system the critical parameters during reaction are temperature of 225°–285°C, amount of water 5–10% and stabilizer content of 1% and time of polymerization.
The DP of Nylon 6 is governed by the following relationship

\[ DP = \frac{(p + r + q)}{(1 + r)} \left(1 - p\right) \]

where

\[ p = 1 - \left\{ \frac{(X + Y)}{(X_0 + Y_0)} \right\} \]

\[ q = \frac{X_{00}}{X_0} \]

and

\[ r = \frac{Y_0}{X_0} \]

Here \( X_0 \) and \( X \) represents the carboxyl group concentration at the start of polymerization and after some time, \( t \), respectively. \( Y_0 \) and \( Y \) represent the amino group concentrations at the initial and final stage, respectively. \( X_{00} \) represents the initial concentration of carboxyl group of a monofunctional acid acting as a stabilizer. Acetic acid is typically used as a stabilizer.

### 2.3.2 Monomer synthesis for nylon 6

**Caprolactam**

There are three important commercial processes for the caprolactam synthesis and all processes start from materials that belong to the group consisting of phenol, benzene, toluene, and cyclohexane. The principal intermediates are cyclohexanone and cyclohexanone oxime for process 1, cyclohexanone oxime for process 2, and cyclohexane carboxylic acid for process 3. The cyclohexanone may be produced by any of the following methods:
1. Catalytic hydrogenation of phenol and subsequent dehydrogenation of the resulting cyclohexanol.
2. One step catalytic hydrogenation of phenol using a palladium on carbon catalyst.
3. Catalytic oxidation (air) of cyclohexane to a cyclohexanol/cyclohexane mixture.
4. Reductive catalytic hydrolysis of cyclohexylamine in one stage operation.

Though process 1 is still practiced, 2 and 3 are the most significant commercial processes. Cyclohexanone oxime may be produced by any of the following methods.

1. Reaction of cyclohexanone with hydroxylamine (process 1).
2. Photonitrosation of cyclohexane with nitrosyl chloride (PNC process) (process 2).

The hydroxylamine used for the reaction with cyclohexanone is obtained by the Raschig process or by catalytic hydrogenation of either nitric oxide or nitric acid.

\[
\text{NH}_4\text{NO}_2 + \text{NH}_4\text{OH} + 2\text{SO}_2 \xrightarrow{5^\circ \text{C}} \text{HON} (\text{SO}_3\text{NH}_4)_2
\]

\[
+ \text{H}_2\text{O}, 100^\circ \text{C} \rightarrow (\text{NH}_3\text{OH})\text{HSO}_4 + (\text{NH}_4)_2\text{SO}_4
\]
(Raschig process)

\[
\text{NO} + 1.5\text{H}_2 + \text{H}_2\text{SO}_4 \rightarrow (\text{NH}_3\text{OH})\text{HSO}_4
\]
(Catalytic hydrogenation of nitric oxide)

The catalytic hydrogenation of nitrate ions involves Pd/C as catalyst in a phosphoric acid ammonium hydrogen phosphate buffering system:

\[
\text{HNO}_3 + 3\text{H}_2 + \text{H}_3\text{PO}_4 + \text{NH}_4\text{H}_2\text{PO}_4 \rightarrow (\text{NH}_3\text{OH})\text{H}_2\text{PO}_4 + \\
\text{NH}_4\text{H}_2\text{PO}_4 + 2\text{H}_2\text{O}
\]

Since the hydroxyl ammonium ion is unstable, the reaction product is directly contacted with cyclohexanone in a toluene solution to effect formation of cyclohexanone oxime:

\[
\text{C}_6\text{H}_{10} + (\text{NH}_3\text{OH})\text{H}_2\text{PO}_4 \rightarrow \text{C}_6\text{H}_{11}\text{ON} + \text{H}_3\text{PO}_4 + \text{H}_2\text{O}
\]

The aqueous H$_3$PO$_4$–NH$_4$H$_2$PO$_4$ solution is reacted with a mixture of NO and NO$_2$ which results in the conversion of any ammonium ions to nitrogen:

\[
2\text{NH}_4^+ + \text{NO} + \text{NO}_2 \rightarrow 2\text{N}_2 + 2\text{H}^+ + 3\text{H}_2\text{O}
\]
Simultaneous introduction of air results in the formation of nitric acid which is recycled in the process.

\[ 2\text{NO} + 1.5\text{O}_2 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_3 \]

An interesting process developed and operated since 1962 by Toray for manufacturing cyclohexanone oxime is the photo-nitrosyl chlorination (PNC) process. This process may be explained by the following mechanism:

\[ \text{NOCl} \xrightarrow{h\nu} \text{NO}^- + \text{Cl}^- \]

\[ \text{C}_6\text{H}_{12} + \text{Cl}^- \rightarrow \text{C}_6\text{H}_{11}^- + \text{HCl} \]

\[ \text{C}_6\text{H}_{11}^- + \text{NO} \xrightarrow{\text{HCl}, <20^\circ\text{C}; h\nu} \text{Cyclohexanone oxime hydrochloride} \]

The nitrosyl chloride for synthesis is obtained by a sequence of reactions that includes:

1. Oxidation of ammonia: \(2\text{NH}_3 + 3\text{O}_3 \rightarrow \text{NO.NO}_2 + 3\text{H}_2\text{O}\)
2. Formation of nitrosyl sulfuric acid: \(2\text{H}_2\text{SO}_4 + \text{NO.NO}_2 \rightarrow 2\text{NOHSO}_4 + \text{H}_2\text{O}\)
3. Sulfuric acid regeneration: \(\text{NOHSO}_4 + \text{HCl} \rightarrow \text{NOCl} + \text{H}_2\text{SO}_4\)

Toluene is the starting material of a commercial process developed by SNIA Viscosa. It involves the following reactions:

\[ \text{Toluene} \xrightarrow{\text{O}_2, 160^\circ\text{C}} \text{C}_6\text{H}_5\text{COOH} \xrightarrow{\text{H}_2, 170^\circ\text{C}} \text{C}_6\text{H}_{11}\text{COOH} \xrightarrow{\text{NOHSO}_4, \text{H}_2\text{SO}_4 \cdot \text{SO}_3, 80^\circ\text{C}} \text{Caprolactam} \cdot \text{H}_2\text{SO}_4 + \text{CO}_2 \]

Reaction of cyclohexane carboxylic acid with nitrosylsulfuric acid in oleum results directly in the formation of caprolactam sulphate by a mechanism that entails simultaneous nitrosation, decarboxylation, and rearrangement of the formed oxime.

2.3.3 Molecular weights suitable for filament production

With appropriate molecular weights, it is possible to produce filaments through melt spinning process. Typically the number average molecular weight used for fiber production process is 18000–20000. Technical grade nylons are made from 24000–26000 number average molecular weight polymers. These high molecular weight nylons are produced by solid state
polymerization from low molecular weight chips obtained through melt polycondensation process [7].

2.4 Thermal properties of nylon fibers

2.4.1 Glass transition temperature

Glass transition temperature (Tg) is associated with amorphous regions and has a significant effect on both processing and physical properties of nylon filaments. The segmental mobility during glass transition in nylon is associated with both flexibility of \(-(CH_2)_n\)- sections, as well as to inter- and intra-molecular hydrogen bonding in the molecular chains. In nylon fibers, the most critical parameter for Tg is moisture regain, which is associated with flexibility introduced by water molecules acting as plasticizer. Amorphous nylon 6 has Tg varying from around -10°C in wet conditions to around 90°C in dry conditions. Tg also depends on number of (CH2) units: as (CH2) unit increases, Tg decreases [8]. In the semi-crystalline state of the filament, glass transition temperature is highly dependent on crystallinity, orientation as well as on moisture regain and the rates of testing. Typical Tg values for nylon filaments are in the range of 40°C to 55°C and increase to around 90°C in dry conditions. Dynamic mechanical methods give higher value of Tg by 20°–30°C, as compared to DSC method.

In the melt spinning process, essentially dry nylon is melted and spun in filament form. Filaments absorb moisture in the various steps of production process, i.e. during quenching, spin finish application and further processing. Thus, Tg dependence on varying moisture conditions during production processes is quite critical. Tg is known to have a non-linear relationship with moisture regain for amorphous nylons [9–12]. There have been several attempts to explain the Tg and moisture regain relationship [10, 13–15]; a reasonable explanation is obtained by three step moisture absorption [12] by nylons. The nature of absorption of molecules [16] is shown in Figure 2.1. In the process of moisture absorption from dry nylon, the initial water molecules form double hydrogen bonds between two carbonyl groups by means of free electron pairs on oxygen atoms and may be assessed as firmly bound. Water molecules are also attached by replacing the hydrogen bond between two carbonyl and amide groups and may be classified as loosely bound molecules. Further absorption can take place by multilayer formation of water molecules. These three different kinds of water molecules affect the Tg in a very different manner and this in turn affects the processing and properties of nylon fibers.

In the case of drawn nylons, some of the observations on Tg vs. regain are still valid. The results on linear expansion coefficients of Nylon 6 both parallel and perpendicular demonstrate this [17]. With increasing moisture
regain from dry state, linear thermal expansion coefficient parallel to the fibre axis ($\alpha_{II}$) increases linearly. However, linear thermal expansion coefficient perpendicular to fibre axis ($\alpha_{\perp}$) is negative up to 2% regain level, increases up to 4–5% of regain level and stabilizes thereafter. This particular variation of $\alpha_{\perp}$ can be explained in terms of varying intermolecular interactions between water and nylon molecule in the above regain range. The continuous increase in $\alpha_{II}$ with regain level shows that there will be corresponding variations in properties.

Modulus of nylon fibre at low temperature has shown that initially, as regain increases from the dry state, modulus increases [18]. This implies that the initial water molecules are tightly bound and have a partial specific volume less than unity, thus making the state very rigid. Variation of torsional rigidity of nylon 6 fibers [19] is a function of regains and observes a maximum rate of change at 3% ratio regain. This corresponds to increased mobility [20] of loosely bound water molecules, as observed from NMR measurements at these regain levels. Stress ageing experiments on nylon 6 filaments indicate that stress induced microcrystal formation takes place during moisture absorption, in the regain range of about 2%.

Choice of moisture regain in the quenching zone in nylon filament production process is quite critical, as the moisture regain is changing rapidly. An RH level of 70% will be helpful, as the filaments will quickly attain the regain levels of ~3% in the highly dynamic filament formation process. Drawing process of nylon 6 is also carried out at around 55% RH corresponding to 3% regain levels. Choice of this level is related with minimum changes in Tg of the fiber with fluctuations in relative humidity in this range.

2.1 Interaction of amide groups with water in nylon.
2.4.2 Melting temperature

Melting of nylon fibers is typical to semi-crystalline polymers, occurring in a range of temperatures. The melting range of nylon 6 is in the temperature range of 215°–228°C, while the nylon 66 melting range is between 250° and 265°C. Melting of nylons is dependent on concentration of amide groups and number of CH₂ groups linking these groups. The variation of melting with number of CH₂ groups per monomer unit is shown in Figure 2.2 [21]. The figure further indicates that the melting also depends on symmetry of the structural units. Heat of fusion of α-crystal form for nylon 6 is 64 cal/g., whereas nylon 66 has a value of 61 cal/g.

2.5 Physical structure of nylon fiber

2.5.1 Steric polarity in nylons

In general in polyamids, steric polarity results from invariable sequences of the CO and NH group [5]. Steric polarity is absent in polyamides of XY type, i.e. nylon 66, but is a distinct characteristic of Z type i.e. nylon 6. In the absence of directionality of XY type nylons, lattice disorders may be readily corrected by movement of chain along the chain direction. As shown in Figure 2.3 [22], steric polarity is absent in nylon 66 but is present in nylon 6. Stable polymorphic modifications i.e. α and γ crystals are possible in nylon 6 due to the directionality effect. Similarly, nylon 4 also has two highly distinguishable crystal forms, i.e. α and γ.
2.5.2 Semi-crystalline structure of nylon fibers

Nylon fibers are semi-crystalline in nature and the crystalline structure is significantly affected by the manner of hydrogen bonding between –CO- and –NH- molecules. Among the two crystal forms of nylon 6, α-crystal is characterized by anti-parallel extended chain arrangement in sheet-like structure, with hydrogen bonds between adjacent chains. The stacking of these sheets is marked by an alternate up and down displacement of about 0.37 nm parallel to chain direction. The crystal form is monoclinic. The crystalline unit cell has eight repeat units [23].

Another crystal modification of nylon 6 is the γ crystal form. Adjacent chains in the γ form are aligned parallel to one another; however these chains are no longer fully extended but have a slight twist. The hydrogen bonding between adjacent chains is at 50° to the chain direction. In both crystal forms, hydrogen bonding is complete between amide and carbonyl groups. The strength of hydrogen bond in the α form is higher than the γ form. Due to differences in the two crystal forms, the density of the α form of 1.23 gm/cc is significantly higher than that of the γ form, having a value of 1.16 gm/cc. The γ form is stable only up to a temperature of around 170°C and gets converted to the more stable α form at higher temperatures.

The filament production process is very sensitive to generation of different crystal forms. At low speed of spinning, the α-crystal form is dominant. At high speed of spinning, the γ-crystal form dominates (e.g. high speed spinning). Subsequent drawing process is easier with the γ form, being more deformable. The drawn and heat stabilized filaments have a dominant α-crystal form. Nylon 6, specifically as-spun filament-form contains a mixture of α, γ and pseudo-hexagonal forms. Rapid quenching of nylon 6 forms a molten state, resulting in bundle-like arrangement of chain segments of parallel and anti-parallel directionality, referred to as a
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converted to the α or γ form depending on subsequent processing of spun filaments. However the α form, being more stable, dominates the stabilized filaments.

Figure 2.4 is a representative equatorial diffraction pattern of nylon 6 samples containing α and γ crystal forms along with amorphous regions. Such analysis can in general be used to obtain fractions of the crystal forms and crystallinity.

Nylon 66 has two very similar crystal forms i.e. α and β. The α-crystal form has an anti-parallel chain in sheet form, very similar to that of the α form in nylon 6. The stacking of hydrogen bonded sheets is characterized by a perpendicular distance of 0.36 nm and a displacement of successive sheets of about 0.5 nm in the chain direction. The α and β forms are differentiated by displacement of sheets in the chain direction being strictly in one direction for the α form and an alternate up and down for the β form. In the case of nylon 66, the α-form is slightly more stable than the β-form. Rapid cooled spun nylon 66 filaments also have a bundle-like arrangement of chains with pseudo-hexagonal arrangements, which get converted to stable crystal forms in subsequent processes of drawing and heat setting.

The actual crystal densities in a given fiber may be 1.19 gm/cc for the α-crystal form and 1.14–1.15 gm/cc for the γ-crystal form. The pseudo-hexagonal structure density is in the range of 1.13 gm/cc and a value of 1.09 gm/cc has been reported for amorphous phase.
Amorphous regions in fibers generally have an imposed orientation, in an otherwise random arrangement of molecular chains and designated as amorphous orientation. This could be thought of as chain extension being highest along the fiber axis. The actual amorphous density of the fibers is higher than 1.09 gm/cc, due to the presence of amorphous orientation.

2.5.3 Morphological structure

A reasonable morphological model for nylon fibers is described in terms of the three phase model with microfibrillar and inter-microfibrillar regions as shown in the Figure 2.5 [25]. It should be noted that a two phase model, consisting of crystalline and amorphous regions in fibers, does not adequately describe the fiber structure. The width of the microfibrils ranges between 60 and 200 Å. The microfibrils in turn consists of crystalline and amorphous regions in a series mode. Industrial grade filaments typically have crystalline and amorphous region lengths of ~60 Å and ~30 Å respectively. These crystalline and amorphous regions, arranged in a regular sequence along the fiber axis, act as a micro-lattice. In fact the regularity is sufficient to give diffraction maxima in the small-angle of x-ray scattering. Longitudinal dimensions of microfibrils are not well defined as is the case with this model type. Microfibrils are surrounded by inter-microfibrillar regions and consist of highly oriented molecular chains and are thus in some degree of pseudo-order [26–28]. Microfibrils in turn form an endless oriented network with branching and fusion more common than endings.

Existence of intra-fibrillar tie molecules has been proposed by Peterlin [29]. A comprehensive review of structural models has been recently published [30]. In the drawing process, nygons deform through micro fiber slippage forming a large number of interfibrillar tie molecules which appear
as a separate phase. Thus the modulus and strength of nylons are dominated by the amount of interfibrillar extended chains and should be considered as a separate phase.

There is a significant interaction between microfibrils in nylons. The extended chains in interfibrillar regions are the strongest and have a profound effect on fiber strength. Thus strength can be increased by shearing off the surface of the microfibrils to make an interfibrillar domain. In the highly oriented state, microfibrils tend to fuse via epitaxial crystallization of extended chain molecules as shown in Figure 2.5.

2.6 Mechanical behavior of nylon fibers

A typical stress-strain curve of nylon tire yarns is shown in Figure 2.6. It is interesting to note that the initial modulus of the nylon is quite low. This is followed by a yield point and a sigmoid type of deformation as shown in the stress-strain curve. This behavior of nylons is unique and is related to combination of flexible – (CH₃) – sections, occasional breakage of hydrogen bonds in amorphous regions and physical linking offered by the presence of crystalline regions. Further, nylons have large extensions to break combined with relatively high strength and thus result in high work of rupture.

2.6.1 Strength

The strength of nylon filaments can be assessed on the basis of the morphological models of these fibers as shown in Figure 2.6. The inter micro-
fibrilar phase is the strongest element of the fiber. On the other hand the microfiber deformation is dominated by the presence of an amorphous region in series with the crystalline regions. On the other hand, the regular chain folding makes the microfibrils relatively weak with strength in the order of 2 gm/denier \([31]\). The amorphous domains are the weakest element and initiation of cracks is expected to start from them. In the case of highly drawn filaments, the microscopic crack \([32]\) dimensions are similar to the microfibril diameter of \(\sim 75 \, \text{Å}\). The number of amorphous domains per unit cross-sectional area is estimated to be \(\sim 2.4 \times 10^{11} \, \text{cm}^{-2}\). Taking the long period as 90 Å, concentration of crack nucleation sites is around \(2.6 \times 10^{17} \, \text{cm}^{-3}\). This value is near to the concentration of free radicals found in the breakage of nylon 6 fiber.

One important conclusion from these observations is that the microcracks arising out of amorphous regions do not grow but become unstable and lead to catastrophic failure. Further, the failure is associated with minimum number of chain scissions. Morphological models \([33]\) for low draw ratio and high draw ratio illustrate these facts as shown in Figure 2.7. Thus the strength depends on: (a) volume function of interfibrillar extended chain dimensions; (b) lateral dimension of microcracks; and (c) distance between the crystallites. The later factors reduce the stress at the tip of the microfibrils.

2.6.2 Modulus

Tensile and bending moduli are important properties of the fibers, affecting the final product properties. For a given fineness, bending modulus is proportional to the tensile modulus. Thus, the tensile modulus values are of significance and a lot of information is available. Crystal modulus values
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(Ec) of select nylon 6 filaments having different draw ratios. For highly drawn samples, λ is much less than 1, which shows a dominant parallel response between crystalline and amorphous regions. Further λ, is approximately equal to φ, indicating a degree of isotropy in parallel and perpendicular directions. On the other hand the sample with the intermediate draw ratio, λ ≈ 1 and φ ≈ 0.57. This corresponds to maximum anisotropy, i.e. coupling along the fiber axis is close to series response and in the perpendicular direction, it is a parallel response. Thus a simple two phase model approach for nylon 6 gives satisfactory explanation for modulus values.

Table 2.2 gives the value of all the relevant parameters i.e. Ef, Ec, Ea, λ, φ for three different nylon 6 filaments having different draw ratios. For highly drawn samples, λ is much less than 1, which shows a dominant parallel response between crystalline and amorphous regions. Further λ, is approximately equal to φ, indicating a degree of isotropy in parallel and perpendicular directions. On the other hand the sample with the intermediate draw ratio, λ ≈ 1 and φ ≈ 0.57. This corresponds to maximum anisotropy, i.e. coupling along the fiber axis is close to series response and in the perpendicular direction, it is a parallel response. Thus a simple two phase model approach for nylon 6 gives satisfactory explanation for modulus values.
2.6.3 Elastic recovery behavior

Nylon fibers have excellent elastic recovery behavior. This unique behavior relates to decrease in entropy on tensile deformation and recovery on stress removal, typical of time dependent rubber elastic effects. During deformation of nyons, the \(-\text{(CH}_2\text{)}-\) sections in amorphous regions are typically mobile above the $\beta$-transition temperatures of $\sim -30^\circ\text{C}$ for nyons. However, regular restrictions are imposed in the mobile sections by hydrogen bonds. These mobile sections may be bigger in the absence of hydrogen bonds, but eventually get restricted as the chain enters the crystalline regions. A chance breaking of hydrogen bonds results in the creep behavior of nyons. On removal of load, the recovery process is due to entropic effects and occurs through breaking and reformation of hydrogen bonds. Typically crystallinity of the order of 50–70% in nylon filaments restricts the overall flow of molecular chains, and insures excellent recovery of nyons. It is worth mentioning that different nyons with varying hydrogen bond density have essentially the same recovery behavior. Interestingly, these observations hold for fairly large deformations of nylon filaments.

2.6.4 Relaxation behavior of nylon 6 and nylon 66 tire cord

Both nylon 6 and nylon 66 have been used in a range of applications interchangeably depending on local availability. For industrial grade applications, differences between nylon 6 and nylon 66 become apparent. Comparative relaxation behavior of these two classes of high tenacity nylon filaments is described, based on dynamic mechanical behavior and morphological parameters [34]. Tan $\delta$ peak for nylon 66 is at $\sim 99^\circ\text{C}$ and for nylon 6 it is at $88^\circ\text{C}$. Due to lower $T_g$, the heat generation builds up faster from room temperature for nylon 6 than that for nylon 66. Nylon 6 also has higher loss peak height than that of nylon 66, whereas nylon 66 has broader loss peak than nylon 6. Nylon 6 also has slightly lower crystallinity than...
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Table 2.3 Comparison of morphological factors for highly drawn nylon 6 and nylon 66 filaments

<table>
<thead>
<tr>
<th></th>
<th>Degree of crystallinity, %</th>
<th>Amorphous orientation</th>
<th>Long period, Å</th>
<th>Size of amorphous domain, Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nylon 6</td>
<td>67</td>
<td>30</td>
<td>86</td>
<td>28.7</td>
</tr>
<tr>
<td>Nylon 66</td>
<td>70</td>
<td>36</td>
<td>91</td>
<td>30.3</td>
</tr>
</tbody>
</table>

that of nylon 66. The higher loss peak along with other factors indicates greater heat generation for nylon 6. Nylon 66 has higher thermal stability due to higher melting temperature as compared to nylon 6. These factors combine together to make nylon 66 filaments superior in demanding industrial applications. For these reasons, applications such as airplane tires are made entirely from nylon 66 cords. These observations are supported by morphology data as given in Table 2.3 and these differences relate to crystallinity, integral breadth in amorphous orientation, long period and size of amorphous domains.

2.6.5 Compressive stresses due to twisted structure

Generally twisted structures are used as nylon tire cords as well as other industrial applications. Twisted structures have large compressive stresses and on tensile deformation, these stresses increase significantly. It is likely that deformation reaches the non-linear limit, as this limit is rather low in compression. On tensile loading of the twisted structure, there is further increase in compressive stresses and thus further decay in properties. For these reasons rope structures with a core of parallel threads and a sheath of braided structures is preferred over a twisted structure for industrial applications using large loads.

2.7 Applications of nylon fibers

Due to good elastic recovery, low initial modulus, excellent abrasion resistance and high resistance to rupture, nylon has been used in a very large number of applications. Nylon fibers are used in both apparel and industrial sectors. Light weight and sheer garments are produced from nylon 6 and nylon 66, where low modulus, high strength and good abrasion resistance are of particular importance. Fabrics made from 15 denier monofilaments are used as ladies’ stockings. They show excellent shape retention due to the elastic recovery behavior of nylon. Setting in steam allows
dimensional stability, permanent pleating while liveliness is also reduced. These fabrics may not require ironing. Socks are mostly made from nylon due to similar reasons. Fabrics made from fine filament are extensively used for sarees. Furs from nylon are also popular due to recovery behavior and long life. Wool is blended with nylon to improve its durability.

In technical textiles, nylon fibers are widely used for a number of applications. Nylon is amongst the best carpet fibers because of resilience and excellent aberration resistance and is very useful under heavy traffic. Nylon bulk continuous filament is used for this purpose. Safety belts in cars, hoses, typewriter ribbons, and light weight canvas for luggage are exclusive domains for nylon filaments. Polyurethane coated nylon fabrics are used for making hot air balloons.

Multifilament nylon yarns find extensive applications for reinforcing rubbers such as tires, with the filament denier ranging from 840 to 1620. These are twisted as multi-strand cord, which improves flex fatigue and with high surface areas for easier bonding with latex. In fact most tires for trucks and airplanes are made from nylon tire cords. Nylon ropes and cordages have strength, durability, and resistance to water. Fishing nets are mostly made from nylon twine due to excellent elastic recovery and high wet strength. Sail cloth made from nylon allows deformation due to wind and recovery on reduction of wind speed, and thus allows optimum advantage of the wind speed.

Strimmer lines used for cutting grass are dominantly made from nylon 6, nylon 66, and nylon 69 monofilaments [35]. These filaments provide the correct balance of flexibility, abrasion resistance and thermal dimensional stability. There is a rapid change in local temperature of these lines and stabilized nylon restricts fusing. A very high diameter uniformity and low shrinkage are required for these filaments. Nylon monofilament of 1.5 mm diameter is used for garden strimmer lines and a diameter between 2.0 to 3.0 mm for agricultural cutters.

Monofilament of polyester and nylon are woven to make conveyor belts for manufacture of paper. The fabric is open mesh with good dimensional stability having a wide width of up to 10 m. Typical dimensions of monofilaments used are in the range of 0.15–0.30 mm. Nylon filaments are in the underside of the conveyor in a nylon/polyester combination to improve wear resistance against the machine frame. Further, nylon filaments do not split longitudinally under compression from machine components. Using small groups of twisted monofilaments also further dissipates the compressive forces and a resilient bare weave is achieved. Umbrella cloth is made from nylon due to diameter swelling behavior. Ribbons for printers, bolting cloths, sutures and toothbrush bristles are further applications of thenylons worth mentioning.
2.8 References


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3.1 Introduction

Polyethylene terephthalate (PET) is a linear polyester consisting of organic compounds of repeating ester groups (–COO–). The usefulness of polyester for fibres was discovered when terephthalate acid was incorporated into the repeat unit of the polymer molecule. The PET textiles made possible the increasingly wider applications of synthetic fibres due to the outstanding physical and chemical characteristics which polyester fibres possess in comparison with natural and synthetic fibres; such as, excellent dimensional stability and sturdiness, a high degree of crease resistance, good bulk elasticity, and warm handle. Further PET characteristics such as great stability of thermoset creases, a high degree of light and heat resistance, and durability of polyester textiles, due to their great fibre strength and elongation, are also very good. Finally, the care of polyester is facilitated by quick drying due to the low water sorption and their relatively good resistance to washing treatments and bacteria and microbes. As a result, mass production of PET fibres enhanced the manufacture of other synthetic fibres.

The use of terephthalic acid with ethylene glycol for the development of PET fibres is introduced as follows. Esterification results from the reaction of carboxylic acid with alcohols in the presence of heat and inorganic acid; such as H$_2$SO$_4$, in accordance with the following scheme:

$$\text{RCOOH} + \text{ROH} \xrightarrow{\text{H}^+} \text{RCOOOR'} + \text{H}_2\text{O}$$

\[\text{R may be alkyl or aryl} \quad \text{R'} \text{ is usually alkyl}\]
During esterification, it is always necessary for a small molecule (e.g., water) to be rapidly removed from the reaction mixture. This is called a condensation reaction. When two functional compounds are used in a condensation reaction, they will produce a condensation polymer. In general, a polymer is formed from joining many molecules by condensation reaction. Almost any condensation reaction can be used to prepare polymers provided that the reaction is performed with monomers having two functional groups. For example, the reaction between a compound with two alcohol groups and a compound with two carboxylic acid groups forms a polymer. The polymers can be classified according to the chemical group linking the monomer units. Polyester is a polymer with repeating units linked by an ester group.

Polyester was first prepared in 1863. Bischoff made polyester from dihydric phenol and dicarboxylic acid in 1902. Fischer prepared polyester from \( p \)-hydroxybenzoic acid in 1908–1910.\(^1\) Carothers\(^2\) dealt with the condensation polymerisation and the structure of polyester. He was the first to produce spinnable polyester of high molecular weight by condensation reaction \( \alpha, \omega \)-diols with alkanedioic acids, or by condensation reaction \( \omega \)-hydroxyalkanoic acids. Moreover, he published a series of papers about polyester in the period 1928 to 1929. Carothers\(^3\) prepared a series of esters from aliphatic diols of formula \( \text{HO(CH}_2\text{)}_y\text{OH} \) and aliphatic dicarboxylic acids of formula \( \text{HOOC(CH}_2\text{)}_x\text{COOH} \). The reaction proceeds as:

\[
\text{HOOC(CH}_2\text{)}_x\text{COOH} + \text{HO(CH}_2\text{)}_y\text{OH} \rightarrow -(\text{OC(CH}_2\text{)}_x\text{COO(CH}_2\text{)}_y\text{O})n^- + 2n \text{H}_2\text{O}
\]

where \( y \) was 2, 3, or 6 and \( x \) was 1, 2, 4, 8. The polyester was produced by alcoholysis; i.e., by heating glycol with oxalate, when alcohol was eliminated. The reaction is,

\[
\text{ROOC–COOR} + \text{HO(CH}_2\text{)}_x\text{OH} \rightarrow -(\text{COO(CH}_2\text{)}_x\text{OOC})n^- + 2n \text{ROH}
\]

where \( \text{R} \) was \( \text{C}_2\text{H}_5 \) or \( \text{C}_3\text{H}_7 \) and \( x \) was 10. Polyester produced by polymer/monomer interconversion such as the lactones of \( \delta \)-hydroxyvaleric acid is shown below.

\[
\begin{align*}
\text{H}_2\text{C} & \quad \text{CH}_2 \\
\text{H}_2\text{C} & \quad \text{CO} \\
\text{CH}_2 & \\
\hline
\text{n} & \rightarrow \quad -(\text{O–CH}_2\text{–CH}_2\text{–CH}_2\text{–CH}_2\text{–CO})_n
\end{align*}
\]
The ability of bifunctional lactones has been discussed by Carothers et al.\textsuperscript{4} It was found that the five-membered ring systems stand out to be very stable. The six-membered ones can be converted into linear polyester and by proper treatment back to the monomeric form. Because of intermolecular reaction or intramolecular reaction, a bifunctional compound such as α-hydroxy acid may yield both diametric cyclic ester and a linear polyester.

![Diagram of a bifunctional compound](image)

Cyclic esters with rings of six or more members can be depolymerised into linear polyester. In the reaction of a bifunctional compound, however, the probability that the chain takes up a configuration allowing the ends of the molecule to react is low. Moreover, such ring configurations will be few in number compared with all possible ones. The probability that the two ends of a long chain will occupy positions adjacent to one another, and hence be in a position to react intramolecularly, varies roughly as (chain length)^{−3/2}. As a result, intramolecular reactions become less probable as the number of atoms in the molecule increases.

The lactones of δ-hydroxyvaleric acid formed from the corresponding ω-hydroxy acid produce a linear polyester as follows:

![Diagram of a linear polyester](image)

In polymer preparation, the compound with six-membered or beyond cyclic lactones is quite readily transformed to polymer. Carothers and Hill pro-
duced a polyester melt (m.p. 75°C) by polycondensation of ethylene glycol and sebacic acid. The polycondensation was carried out as part of a molecular distillation or simply by passing nitrogen through the condensation melt until a fibre-forming polyester resulted. The filaments were then processed by cold drawing. They also produced other fibre-forming polyesters; such as the reaction products from propylene glycol and hexadecane dicarboxylic (octadecanedivinc) acid (m.p. 75°C), propylene glycol and \(\omega\)-hydroxydecanoic acid (m.p. 65°C), and propylene glycol and \(\omega\)-hydroxypentadecanoic acid (m.p. 95°C).

In 1931, Carothers produced a polyamide ‘66’ or so-called ‘nylon’ from adipic acid and hexamethylene diamine. Further, Carothers and his collaborators found very large differences in melting points between polyesters and the corresponding polyamides. Polyester melt (m.p. 57°C) resulted from reaction of adipic acid with ethylene glycol, while a polyamide melt (m.p. 278°C) was produced by the reaction of adipic acid with tetramethylene diamine. Also, a polyester melt (m.p. 75°C) was formed by the reaction of sebacic acid with ethylene glycol, while a polyamide melt (m.p. 254°C) was produced by the reaction of sebacic acid with ethylene diamine. Because of the thermal stability and the fact that the melting points of the polyesters produced from aliphatic dicarboxylic acids and glycols were considerably lower than those of the corresponding polyamides, Carothers and his collaborators turned their attention to polyamides to exploit their discovery commercially.

In 1941, Schlack implemented terephthalic acid for the development polyester fibres in Germany at a branch of Agfa Wolfen. He focused more on producing polyester from terephthalic acid and 1, 4-butanediol (m.p. 218°C). At the same time, Whinfield and Dickson applied it at Calico Printers’ Association in Great Britain, while they preferred using a combination of terephthalic acid and ethylene glycol to produce polyester melt (m.p. 268°C). Both polyesters are crystalline and quite adequate for forming fibres. Nevertheless, from the point of view of economy and application to textile usage, the combination selected by Whinfield and Dickson was mostly preferred so that, industrially, the polyester fibres produced from terephthalic acid and ethylene glycol have maintained their competitive importance.

The relationship between the work of Carothers, Schlack, and Whinfield and Dickson is shown in the following diagrammatic representation:

\[
\text{Dicarboxylic} \quad \text{Glycol} \\
\text{HO–(R_2–OOC–R_1–COO)_nR_2–OH} \\
\text{Polyester}
\]

The relationship between the work of Carothers, Schlack, and Whinfield and Dickson is shown in the following diagrammatic representation:
In 1947, mass production, based on the technological development of polyester fibres originated by Whinfield and Dickson, was conducted in Great Britain by Imperial Chemical Industries Ltd (ICI) and in America by the Du Pont de Nemours Company after both companies had acquired the patent rights from the Calico Printers’ Association Ltd, in Manchester. ICI marketed polyester fibre as ‘Terylene’, whilst the fibre produced by Du Pont was called ‘Dacron’. Then, the patent rights covering the polyester fibres developed by the Calico Printers’ Association Ltd and ICI were made available to other firms in France, Italy, etc.

Fibre-forming polyester must possess the necessary properties (melting point, average molecular weight, dyeability, resistance to chemicals, etc.) so that it can be utilized commercially. Aliphatic polyester has low melting temperature; examples are:

\[
\text{HO(CH}_2\text{)}_9\text{COOH} \rightarrow \text{(+O(CH}_2\text{)}_9\text{CO)}_n + \text{H}_2\text{O}
\]

Properties of this polyester: crystallization, forming fibre, can be drawn at room temperature, and melting temperature is 76°C, and

\[
\text{HO(CH}_2\text{)}_{10}\text{OH} + \text{HOOC(CH}_2\text{)}_4\text{COOH} \rightarrow \text{(-O(CH}_2\text{)}_{10}\text{OOC(CH}_2\text{)}_4\text{CO)}_n^- + \text{H}_2\text{O}
\]

Properties of this polyester: crystallization, forming fibre, can be drawn in room temperature, and melting temperature is 80°C.

Aromatic polyester has high melting temperature. The reaction is

\[
\text{HO(CH}_2\text{)}_2\text{OH} + \text{HOOC(O(CH}_2\text{)}_2\text{COOH} \rightarrow \text{(-O(CH}_2\text{)}_2\text{OOC(O(CH}_2\text{)}_2\text{CO)}_n - + \text{H}_2\text{O}
\]

Properties of this polyester: rigid, crystallization, forming fibre, can be drawn in room temperature, and melting temperature is 264°C.
The clothing industry requires polyester of higher strength, greater crystallisation, higher resistance to light, etc. Thus, aromatic polyester of higher melting temperature is chosen for fibre production. In future, degradable fibres or green polymer is of great importance. Polyester has more thermal hydrolytic degradation, more alkaline degradation, and thermal oxidative degradation. Polyester is a chemical synthetic degradable polymeric material. In particular, the fibre-forming polyester with low melting temperature is the best material. Therefore, the aliphatic polyester may be selected for producing green polymer and degradable fibres.

### 3.2 Raw materials of polyester fibres in industry

The fibre-forming polyester may be obtained from dicarboxylic acids with diols, hydroxyl acids, or lactones. Commercially, aromatic polyester is applied using ethylene glycol (EG) and dimethyl terephthalate (DMT), or ethylene glycol and terephthalic acid (TPA) to produce polyethylene terephthalate (PET).

Before 1970, polyester was exclusively produced on a commercial scale from ethylene glycol and dimethyl terephthalate. After 1970, Mobil Co. and Amoco Co. were able to successfully develop technologies to produce pure terephthalic acid. Polyester then started to be produced from ethylene glycol with terephthalic acid. The refining processes for crude oil into ethylene, toluene and xylene are shown in Fig. 3.1.

![Diagram of refining processes for crude oil](image)
The producing processes of ethylene glycol, dimethyl terephthalate, and terephthalic acid are discussed in the following sections.

3.2.1 Production of ethylene glycol

Two methods have achieved importance for the manufacture of ethylene glycol including the ethylene oxide method and the ethylene chlorohydrin method.

*Ethylene oxide method*

Ethylene passes through oxidation processes to produce ethylene oxide which is then hydrated to form ethylene glycol. This method is achieved by:

(a) Producing ethylene oxide from ethylene

Direct oxidation and chlorine processes are of great importance for the technical manufacturing of ethylene oxide.

(i) Direct oxidation process

This process involves direct oxidation of ethylene to ethylene oxide according to the following reaction:

\[ \text{CH}_2=\text{CH}_2 + \frac{1}{2} \text{O}_2 \xrightarrow{\text{Oxidation, 150~300}^\circ\text{C}} \text{H}_2\text{C} \xrightarrow{\text{Catalysts}} \text{CH}_2\text{O} \]

The reaction uses metal catalysts (e.g., silver oxide) and the reaction temperature is approximately 250°C.

(ii) Chlorine process

The chlorine process is carried out in two stages. In the first stage, ethylene with hypochlorous acid is transformed into ethylene chlorohydrin.

\[ \text{CH}_2 = \text{CH}_2 + \text{HOCl} \xrightarrow{\text{Cl}_2, \text{H}_2\text{O}} \text{HOCH}_2\text{CH}_2\text{Cl} \]

In the second stage, ethylene chlorohydrin along with calcium hydroxide produces ethylene glycol.

\[ 2\text{HOCH}_2\text{CH}_2\text{Cl} \xrightarrow{\text{Ca(OH)}_2} 2\text{CH}_2 = \text{CH}_2 + \text{CaCl}_2 + 2\text{H}_2\text{O} \]
(b) Production of ethylene glycol from ethylene oxide

Ethylene oxide may react with water and be hydrated to ethylene glycol in accordance with the following reaction:

\[
\text{H}_2\text{C} = \text{CH}_2 + \text{H}_2\text{O} \xrightarrow{\text{Hydrogenation}} \text{HOCH}_2 = \text{CH}_2\text{OH}
\]

The reaction proceeds with the aid of acid catalysts, while the reaction temperature ranges between 60 to 100°C.

**Ethylene chlorohydrin method**

Ethylene glycol is produced by alkaline saponification of ethylene chlorohydrin with caustic soda solution, or of ethylene chloride and soda according to the following formulae:

\[
\text{ClCH}_2 = \text{CH}_2\text{OH} + \text{H}_2\text{O} \xrightarrow{\text{Caustic soda}} \text{HOCH}_2 = \text{CH}_2\text{OH} + \text{HCl}
\]

\[
\text{ClCH}_2 = \text{CH}_2\text{Cl} + 2\text{H}_2\text{O} \xrightarrow{\text{Soda}} \text{HOCH}_2 - \text{CH}_2\text{OH} + 2\text{HCl}
\]

However, this method is one of the older procedures and thus it was abandoned. Ethylene glycol was purified to produce polyester fibres with the following properties:

- Boiling point: 195–198°C.
- Density: 1.110–1.112 (at 20°C).
- OH number: greater than 1750.
- Water content: less than 0.1%.
- Ester interchange value: greater than 90.

### 3.2.2 Manufacturing terephthalic acid and dimethyl terephthalate

Benzene, toluene and xylene are the primary raw materials in manufacturing terephthalic acid and dimethyl terephthalate. The flow chart for the manufacturing processes of TPA and DMT is depicted in Fig. 3.2.

**Benzene material**

The following reaction system shows the conversion of benzene to terephthalic acid:
3.2 Flow chart for TPA and DMT.
Toluene material

Four major processes of manufacturing terephthalic acid and dimethyl terephthalate from toluene including:

1. Synthesis via $p$-tolualdehyde
   The synthesis produces terephthalic acid and dimethyl terephthalate according to the following reaction patterns:

   ![Reaction diagram 1](image1)

   (Toluene) (p-tolualdehyde)

   or

   ![Reaction diagram 2](image2)

   (Dimethyl terephthalate)

2. Synthesis via $\omega, \omega'$-dichloro-$p$-xylene
   The terephthalic acid or dimethyl terephthalate is obtained by three established processes according to the following formulae:

   ![Reaction diagram 3](image3)

   $(\omega$-chloro-xylene) (4-methylbenzyl alcohol)

   or
3. Synthesis via p-tolualide

The manufacture of terephthalic acid based on tolune proceeds as follows:

4. Synthesis via potassium benzoate

The synthesis for the production of terephthalic acid is based on the following reactions:\textsuperscript{11}
Material from \( p \)-xylene

Conversion of \( p \)-xylene mainly occurs by oxidation. Terephthalic acid and dimethyl terephthalate are produced from \( p \)-xylene according to the following formula:

\[
\begin{align*}
\text{CH}_3 & \quad \text{Oxidation} & \quad \text{COOCH}_3 \\
\text{CH}_3\text{COOH} & \quad \text{COOH} \\
\end{align*}
\]

The oxidation of \( p \)-xylene may be carried out in two ways:

(a) One-step oxidation process, which includes catalytic oxidation,\(^{12}\) and the Willgerodt reaction,\(^ {13}\) etc.

(b) Two-step oxidation process,\(^ {14}\) which leads directly to dimethyl terephthalate by the following reaction:
Currently, the one-step oxidation of \( p \)-xylene is commonly adopted. Terephthalic acid and dimethyl terephthalate are purified and then used as raw materials for polyester fibre production.

### 3.3 Polymerisation process of polyester fibres in industry

In industrial practice, the polymerisation processes of aromatic polyethylene terephthalate are produced via a two-step process. The first step is esterification and preliminary condensation, while the second is polycondensation or melt polymerization. The flow chart for polyethylene terephthalate (PET) process is depicted in Fig. 3.3.

#### 3.3.1 Esterification and preliminary condensation

Esterification and preliminary condensation proceed in two paths: (1) transesterification and preliminary condensation and (2) direct esterification.
tion and preliminary condensation. In the first path, the DMT is transformed, with the assistance of one or more catalysts and EG at temperatures ranging from 150 to 200°C, into BHET (bis-(2-hydroxyethyl) terephthalate) (also called diglycol terephthalate, DGT) with the elimination of methanol. The reactions of first path proceed as follows:

**Transesterification**

\[
\begin{align*}
\text{H}_2\text{COOC-} & \text{COOCH}_3 + 2\text{HOCH}_2\text{CH}_2\text{OH} \\
\text{(DMT)} & \quad \text{(EG)} \\
\rightarrow & \text{HOCH}_2\text{CH}_2\text{OOC-} \text{COOCH}_2\text{CH}_2\text{OH} + \text{CH}_3\text{OH} \\
& + \text{DMT} + \text{EG} + \text{catalysts}
\end{align*}
\]

**Preliminary condensation**

\[
\begin{align*}
\text{H}_2\text{COOC-} & \text{COOCH}_3 + \text{HOCH}_2\text{CH}_2\text{OH} \\
\text{(DMT)} & \quad \text{(EG)} \\
\rightarrow & \text{HOCH}_2\text{CH}_2\text{OOC-} \text{CO-}x\text{OCH}_2\text{CH}_2\text{OH} \\
& (\text{BHET}) \quad x=3\sim5
\end{align*}
\]

The reaction conditions of transesterification and preliminary condensation are:16

- Catalysts: metal oxides (PbO, MgO, Sb\textsubscript{2}O\textsubscript{3}, etc.), metal acetates (Co, Mn, Zn, etc.), or mixtures of oxides and acetates metals (CH\textsubscript{3}COO\textsubscript{2}CO-Sb\textsubscript{2}O\textsubscript{3}, (CH\textsubscript{3}COO\textsubscript{2})\textsubscript{2}CO-PbO, etc.).
- Concentration of catalyst: 0.05% (to the amount of DMT).
- Monomer ingredient for DMT: EG = 1: (2–2.5) (mole ratio).
- Reaction temperature: falls in the range of 150–200°C (gradually increases).
- Reaction time: 3–6 hours.

In the second path (direct esterification and preliminary condensation), terephthalic acid (TPA) is converted, with EG and the aid of one or more catalysts at temperatures ranging from 250 to 280°C, into BHET with the elimination of water. The reaction is explained in the following scheme:
Direct esterification

\[
\text{HOOC-COOH} + 2 \text{HOCH}_2\text{CH}_2\text{OH} \xrightarrow{\text{catalysts}} \text{HOCH}_2\text{CH}_2\text{OOCCOOCH}_2\text{CH}_2\text{OH} + \text{H}_2\text{O}
\]

+ TPA + EG + catalyate

Preliminary condensation

\[
\text{HOOC-COOH} + \text{HOCH}_2\text{CH}_2\text{OH} \xrightarrow{\text{catalysts}} 
\]

\[
\text{HOCH}_2\text{CH}_2\text{OOCCO}_x\text{OCH}_2\text{CH}_2\text{CH}_2\text{OH} \quad (\text{BHET}) \quad x=3\sim5
\]

The reaction conditions of direct esterification and preliminary condensation are:

- Catalysts: metal oxides (Sb\textsubscript{2}O\textsubscript{3}, PbO, etc.).
- The concentration of catalysts: should be less than 0.05% (to the amount of TPA).
- Monomer ingredient for TPA: EG = 1: (1.1–2) (mole ratio).
- Reaction temperature: 250–290°C (increases gradually).
- Reaction time: 3–6 hours.

3.3.2 Polycondensation

This step starts by the BHET produced by esterification and preliminary condensation. Then, by condensation reaction, the intermolecular BHET releases EG and produces PET. The reaction is:

\[
\text{HOCH}_2\text{CH}_2\text{OOCCO}_x\text{OCH}_2\text{CH}_2\text{OH} + \text{EG} + \text{TPA(or DMT)} \xrightarrow{\text{X=3\sim5}} 
\]

\[
\text{HOOC-COOCH}_2\text{CH}_2\text{OH} + \text{HOCH}_2\text{CH}_2\text{OH} \quad (\text{PET}) \quad m \approx 100 \quad (\text{EG})
\]
The reaction conditions of polycondensation are:

- Catalysts: metal acetates (e.g., Co, Mn, Zn, Mg, Pb, Cd, etc.) or mixtures of oxides and acetate metals; such as \((\text{CH}_3\text{COO})_2\text{Mn} + \text{Sb}_2\text{O}_3\), \((\text{CH}_3\text{COO})_2\text{Co} + \text{Sb}_2\text{O}_3\), etc.
- The concentration of catalysts: 0.02–0.03% (to the amount of TPA or DMT).
- The thermal stabilizer of PET: phosphor compounds.
- The concentration of the thermal stabilizer of PET: 0.015–0.03% (to the amount of TPA or DMT).
- Nitrogen under pressure is needed to place BHET into the condensation vessel.
- Reaction pressure: below 1 mm/Hg.
- Reaction time: 4–6 hours.

3.4 Fundamental principles and types of melt spinning process

The spinning process of fibre-forming polyester converted to fibre passes into three stages. In the first stage, a solid fibre-forming polyester is produced from polymerisation, which is called polymer chips. Polyester chips are converted to dope by heat, or by dissolving in some solvent.

\[
\text{Polymer chips} \xrightarrow{\text{Conversion}} \text{Dope (viscoelastic fluid, or spinning fluid)}
\]

In the second stage, dope fluid is extruded through a spinneret and then converted to viscoelastic filament (also known as the originally extruded filament). The viscoelastic filament is solidified into solid filament fibre (usually called filament), which may be formed by extending the viscoelastic filament.

\[
\text{Dope} \xrightarrow{\text{Spinning}} \text{Viscoelastic filament} \xrightarrow{\text{Solidification}} \text{Solid filament fibre}
\]

Solidification may be obtained by cooling,\(^{17,18}\) by extruding the viscoelastic filament submerged in a coagulation bath for removal of the solvent,\(^{19}\) or by extrusion into the heating chamber for evaporation of the solvent.\(^{20}\) From both stages, there are three methods of the spinning process:
1. Melt spinning:

Polymer chips $\xrightarrow{\text{Conversion by heat}}$ Dope $\xrightarrow{\text{Spinning}}$

Viscoelastic filament $\xrightarrow{\text{Solidification by cooling}}$

Solid filament fibre

2. Wet spinning:

Polymer chips $\xrightarrow{\text{Conversion by dissolving in solvent}}$ Dope $\xrightarrow{\text{Spinning}}$

Viscoelastic filament

Solidification by submerging in coagulation bath

Solid filament fibre

3. Dry spinning$^{21,22}$:

Polymer chips $\xrightarrow{\text{Conversion by dissolving in solvent}}$ Dope $\xrightarrow{\text{Spinning}}$

Viscoelastic filament

Solidification by curing in the heating chamber

Solid filament fibre

If the polymer chips are thermoplastic polymers (i.e., PET, polyamide, polyurethane, polypropylene, etc.), melt spinning, wet spinning or dry spinning can be used. Usually, melt spinning is used. But if polymer chips are thermosetting polymers (i.e., polyacrylonitrile, viscose rayon, cellulose acetate, etc.), wet spinning or dry spinning can be used.

In the third stage, when the viscoelastic filament is solidified into solid filament fibre, it is extended by the drawing process. The drawing process decreases the diameter, and the orientation of solid filament fibre could increase, while it improves the degree of crystallinity.

3.4.1 Fundamental principles of melt spinning process

PET belongs to the thermoplastic polymer and so the commercial PET fibre usually uses melt spinning. The schematic melt spinning process for PET fibre is shown in Fig. 3.4. PET fibres are formed by extrusion of the molten polymer. Apparently, this is a simple process. A supply of molten PET (D) is pumped at a constant rate and under very high pressure through small holes in the spinning jet (E). The viscoelastic filaments are extruded to emerge vertically downwards from the face of the spinning jet and, on cooling (F), solidify, and wind onto the bobbins.
A. Hopper fed with PET chips
B. Spinning vessel
C. Heated grid
D. Dope (molten PET)
E. Spinning jet (spinneret)
F. Cold air
G. Input feed rollers
H. Deflector
I. Output feed roller

3.4 Schematic melt spinning process for PET fibre.

3.5 The profile model of spinning jet.

In Fig. 3.4, the profile model of dope fluid (viscoelastic fluid) flows through the spinning jet (E), which is described in Fig. 3.5.

In the inlet zone (c), the dope fluid is contracted, and then the polymer chain in the dope fluid produces elastic deformation. In the entire extended zone (d), the velocity of viscoelastic fluid is a steady flow. The contraction phenomenon of viscoelastic fluid in the jet issuing zone (e) is related to the different velocity profiles inside and outside the spinneret hole tube. The fluid jet expands outside the spinneret hole tube and its maximum radius is often called die swell. The profile of total energy transformation of the dope fluid flowing through spinneret hole tube is illustrated in Fig. 3.6.

Dope fluid (viscoelastic fluid) is extruded through the spinneret, and then converted to viscoelastic filament. The viscoelastic filament is transformed into solid filament fibres by cooling, where the viscoelastic filament
will be extended and the diameter of the viscoelastic filament will be reduced. The entire deformation process of the viscoelastic filament by elongation strain can be divided into three sections as shown in Fig. 3.7:

(a) Die swell section with negative parallel (elongation) velocity gradient.
(b) Elongation section (or deformation section) with positive parallel (elongation) velocity gradient.
(c) Solid motion section (running solid filament fibre section) with zero parallel (elongation) velocity gradient or without deformation.
The diameter of the viscoelastic filament is maximum at the intersection of the die swell and elongation sections. The distance is usually less than 10 mm from the spinneret. The distance of the elongation section is usually 50–150 cm from the spinneret. The viscoelastic filament (originally extruded filament) on the spinning axial line is then solidified by cooling. During solidification there are two temperature gradients. The first temperature gradient is along the spinning axial line, while the other is radial which expresses the radial differential temperature from the outer layers to the inner layers of viscoelastic filament. Although the range is only a few degrees celsius, the physical properties of viscoelastic filament may be significantly affected. Fig. 3.8 displays the influence of physical properties on the viscoelastic filament.²⁴

The following results are obtained from Fig. 3.8 regarding:

(a) Temperature: the outer layers are lower, while the inner layers are higher.
(b) Crystallisation rate: the outer layers are fast, while the inner layers are slow.
(c) Viscosity: the outer layers are high; however, the inner layers are low.
(d) Axial spinning velocity: uniform velocity on radial.
(e) Extensional force: the outer layers are high, whereas the inner layers are low.

Drawing of PET viscoelastic filaments or filament (solid filament fibres) may be performed, depending on the fibre type, under cold or hot...
conditions, besides it has an additional effect of making the filament both narrower and longer. The drawing process is usually performed at a temperature that must be higher than the secondary transition temperature ($T_g$) of PET fibres. The drawing process is achieved by stretching the viscoelastic filament between two rollers (input feed roller and output feed roller), called godet rolls, with the output feed roller rotating faster. It also may be performed immediately after spinning of viscoelastic filament or during subsequent processing as part of yarn texturing and/or spinning.

The industrial drawing depicted in Fig. 3.9 is a continuous operation. In this process, the PET filament is supplied at the input feed roller with a constant feed velocity, $V_{in}$, and is taken-up by the output feed roller running with a constant drawing velocity, $V_{out}$, equalling $R \times V_{in}$, where $R$ is the nominal draw ratio. The actual draw ratio is less than $R$ because the drawing filament shrinks when tension is removed.

The stress-strain curve (S-S curve) of the PET viscoelastic filament during the drawing process is shown in Fig. 3.10. From this figure, it is
noted that the deformation in the OA region is uniform and is usually reversible. The slope of S-S curve (called modulus = σ/ε) is constant. The deformation is called elastic deformation. In the AB region, the modulus decreases when the stress is increased. The drawing exhibits a non-uniform deformation. The largest stress corresponds to point B which is often called the yield point. In the BC region, the deformation is unstable, where local attenuation results in a ‘necking effect’ concentration of stress. The strain is called strain-softening. The CD region is called the necking effect region. The deformation is concentrated in one or more necks, which gradually extends over the entire filament, while the diameter of the neck and undrawn parts remains steady. The deformation in this region is irreversible. In the DE region, the deformation is uniform and is usually irreversible. The strain increases as the stress increases. The drawing exhibits a uniform deformation. Point E is called the breaking point. Finally, the OF and OG are called the breaking strain and the breaking stress, respectively.

Crystalline and amorphous arrangements of PET chain exist within newly formed filaments. These PET chain molecules are possibly oriented to make them more parallel to the walls of the filament by stretching the filament before it is completely solidified. This results in more crystalline and stronger filaments. However, not all filaments are drawn to the maximum amount possible, because when a filament fibre reaches its maximum length, the extensibility of filament fibres (or filament yarn) decreases. Therefore, filament yarn can be divided into two types based on the degree of drawing (called the draw ratio). The filaments that have not been fully drawn are called partially oriented yarns (POY), while those that have been fully drawn are called fully oriented yarns (FOY).
3.4.2 Fundamental types of melt spinning processes

There are two procedures for the production process of PET fibre. The first procedure is the production process for PET, while the second one is the melt spinning process for PET fibres.

Production processes for polyethylene terephthalate

There are two production processes for producing PET including batch process and continuous process. These two processes are illustrated as follows:

1. Batch process, which proceeds as follows:

   - **PTA** → **Slurry tank** → **Esterification vessel** → **Filter** → **Vacuum polycondensation vessel** → **Polyethylene terephthalate (PET)**
   - **EG** → **Catalysts**
   - **Catalysts and thermal stabiliser**

2. Continuous process, which is conducted as shown below:

   - **PTA** → **Slurry tank** → **First esterification vessel** → **Second esterification vessel** → **First vacuum preliminary polycondensation vessel** → **Catalysts and thermal stabiliser**
   - **EG** → **Catalysts**
   - **TiO₂**

   - **Second vacuum preliminary polycondensation vessel** → **Filter** → **Vacuum primary polycondensation vessel**
   - **PET** → **Filter** → **PET Direct melt-spinning**
   - **Chips granulator** → **PET chips** → **Melt spinning**
Melt spinning process for polyethylene terephthalate fibres

PET staple fibres and PET filament fibres are produced by melt spinning processes as follows:

1. PET staple fibres production

   - PET chips
   - PET direct
   - Melt-spinning
   - Cooling
   - Collection of fibres
   - Oil immersion
   - Steam heating
   - Extended heating
   - Crimping
   - Heat-setting
   - Cutting (staple fibres)
   - Baling

2. PET filament fibres production

   - PET chips
   - PET direct
   - Melt-spinning
   - Cooling
   - Cold drawing
   - Oiling
   - Winding
   - Bobbin or cake

3.5 Heat setting and textured yarn of filament

3.5.1 Heat setting

In the melt spinning process, the time of fibre-forming is usually short. This results in different relaxing states of polymer chain in the PET fibre and causes non-uniform internal stresses within it. In effect, many crystal defects appear. Moreover, such an unstable structure would cause fibre shrinkage that makes any further finishing processes so difficult. Thus, in order to achieve a stable fibre structure, heat setting should be controlled before the fibres are used.

Heat setting determines the morphology and dimensional stability of thermoplastic fibres. It includes three fundamental factors: temperature,
time and speed. In practice, temperature should be set greater than the secondary transition temperature ($T_g$) and below the melting temperature ($T_m$). Heat setting causes the movement of polymer chains in fibres, which releases the internal stress, finally producing a complete and stable structure of fibres as shown in Fig. 3.11. Dismore and Statton²⁶ show that the polymer chain would change into a fold when the heat setting temperature is increased.

The heat-setting process passes through three stages. The first stage is called the fibre structure loosing stage in which, due to heating, the fibre temperature is higher than $T_g$. However, the interaction force between the PET chain in fibres is weakened in a few seconds. The second stage is an actual heat-setting stage in which the binding energy between the PET chain spontaneously increases, while the PET chain produces heat-vibration. Because of the fibre structure loosing stage and the heat-vibration phenomenon, a new binding force between PET chains (in fibre) is formed. The completion time of this stage is several times that of the first stage. Finally, in the third stage, the binding force and the structure of the PET chain are fixed when the fibre temperature is kept below $T_g$. In practice, this stage can be completed in a few seconds.

### 3.5.2 False-twist process

In 1954, the first machine for false-twist was invented. This machine can produce a filament yarn with stretchability. The fundamental principle of the false-twist process is to change the straight multi-filaments into a crimp filament. In this process, the straight multi-filament is placed at a system above $T_g$. Then, it is twisted with force to re-arrange the internal molecule and form a perpetual deformation. The main characteristic of the false-twist process is to rotate the middle of the multi-filament so that the twist numbers at both sides of the rotation point are equal in each direction²⁷ as shown in Fig. 3.12(a).
The false-twist process shown in Fig. 3.12(b) sets a heater in the twist area while the crimp of the filament is fixed. When the multi-filament is twisted through the pin (or spindle) into the re-twist area, the twist number of the multi-filament becomes zero while each monofilament holds the crimp shape.

The false-twist process includes twist, heat setting and re-twists. Generally, textured yarn is produced by the false-twist process. Two types of false-twist process exist. The first is pin-type spindle twisting, while the other is friction twisting. The mechanisms of these two types are illustrated in Fig. 3.13.\(^\text{28}\)
3.6 High speed spinning and novel spinning

3.6.1 High speed spinning of polyester filaments

The first concept of high speed spinning was developed by DuPont’s US patent in 1952. Along with the development of industrial technology, the high speed take-up machine for above 6000 m/min was developed in 1980. High speed spinning was then commercialized. Due to the continuous development of mechanism and electronic control systems, it is expected that the spinning speed will reach 8000 m/min during the twenty-first century. Meanwhile the spinning speed for research in the laboratory is over 12000 m/min.

Fundamental principles of high speed spinning

High speed spinning was developed based on the conventional spinning method of 1500~2500 m/min speed. This process combines the spinning and drawing processes, which are performed at a speed ranging between 3500~4000 m/min, in the same machine. The take-up of the fibre takes place with a winding speed of 5000~6500 m/min.

The key point of high speed spinning is the orientation of molecules, which is influenced by several spinning parameters; especially spinning tension. Spinning tension is related to the spinning speed, the raw material property, and the processing conditions; including process temperature, the condition of quenching, and the spin-draw ratio of spinneret.

The polymer melt deformation in the high speed spinning process is shown in Fig. 3.14. It is noted that the fibre forming structure is divided into three zones: (I) flow-deformation, (II) oriented-crystallisation, and (III) plastic-deformation. When the polymer melt passes through the spinning jet and emerges from the spin holes into the air, it is normally solidified in a distance of 30~130 cm from the surface of spinneret (depends on the spinning condition). Therefore, the structure of the fibre will be changed. The basic parameters of fibre structure are birefringence, crystallisation, and crystal size. Normally, birefringence and crystallisation of partially oriented yarn are increased by raising the spinning speed of the fibre. In effect, tenacity is increased, while the elongation of the fibre is reduced.

High speed spinning process

Manufacturers prefer a process that produces a full oriented yarn with one single machine. Therefore, the high speed spinning process was developed in various methods.
It is well known that the basic process of melt spinning includes: the melting condition of polymer, the through-put of melt from the spinneret, the thinning process of the polymer melt (viscoelastic filament), the solidification of the viscoelastic filament, and the oil pick-up and fibre take-up procedure.

The high speed spinning process is improved from the conventional melt spinning technology by increasing the spinning speed to 3500 m/min, adding the drawing rollers to three or more heated godets, and raising up the winding speed to 5000–6500 m/min. High speed spinning process is depicted in Fig. 3.15.

In Fig. 3.15, the yarn path goes through rollers 1 and 2, where the speed of roller 2 is greater than 3500 m/min. The drawing process is then performed with different operation speeds of rollers 2 and 3, where the speed of roller 3 could be raised to 5000–6500 m/min. Finally, the yarn is taken-up by a high speed winding machine so that a full oriented yarn is obtained.\(^{33}\)

When using the high speed spinning process, the production capacity may be increased by more than 20%. In return, the production cost would be reduced effectively. However, high spinning speed causes higher winding tension of yarn and, consequently, increases the instability of the spin-line.

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As a result, an optimal oil application and effective performance of yarn guides are critical for the success of this process.

3.6.2 Novel spinning of polyester fibres

Melt blown process

Melt blown process, shown in Fig. 3.16, was initially developed by Naval Research Lab, USA in 1951.\textsuperscript{30} Then, it was classified as melt spun nonwoven process.\textsuperscript{34,35} By this process, almost all the thermoplastic polymer can
be produced with micro-fibre. The diameter of single fibre from the melt blown process is 1 micron or less, but the variation of the fibre size is relatively high. The tensile properties of melt blown nonwoven for machine direction and cross direction show an obvious uneven behaviour. Since melt spinning has an advantage for mass production, researchers focus on the improvements for the melt blown process. For example, the design of a special spinning device and spin pack to reduce the fibre size to sub-microns, where it is expected to achieve a nano-scale fibre with a cost below $US10; Nanoval Co. from Germany was authorised by Neumag Co. to develop an air impingement method to make ultra-fine fibres; and Hills Co. of USA design the thin distribution plates to enlarge the spin holes per unit area of spin pack and thus the through-put per hole will be reduced and consequently a finer fibre can be attained.\textsuperscript{36} In addition, the alternation of L/D for spin hole may help in reducing the melt blown fibre dimension. The ideal diameter of the spin hole is around 0.10–0.15 mm, while the optimum melt flow index of the polymer is controlled at 1000.

\textit{Split conjugate spinning process}

It is very hard to produce a micro-fibre with the denier per filament below 0.3 denier by a conventional single component method.\textsuperscript{37,38} Before 1960, there was no commercialised technology for micro-fibre. In 1970, Toray Co. in Japan started its promotion of micro-fibre products. The split conjugated bi-component fibre is an approach for producing micro-fibres, which includes two incompatible polymers such as PET and PA6, using the conjugate spinning method\textsuperscript{39,40} to spin to fibre. These two polymers are distributed similarly to the petal segments type on fibre cross-section. Each conjugated filament is then divided into several finer fibres by means of a mechanical or a chemical method. It can possibly be split during the fabrication and dyeing process into 12, 16, 24 or 32 segments, and so on, depending on the design of the spin pack. This method is called the split conjugated micro-fibres. After splitting the denier of fine fibre may reach 0.03–0.07 denier. The end product using this micro-fibre is suitable for several products including high quality wipers and high density fabrics for apparel and home textiles. Figs. 3.17 and 3.18 display the schemes of split conjugate fibre for (24 segment-pie) and spin pack for split conjugate fibre, respectively.

The purpose of split conjugated micro-fibre is to get a fine fibre by the splitting of the interface of two polymers. In this process, the appropriate design of conjugate configuration is very important and the viscosity of the independent polymer is critical. If there is a huge difference of flow behaviour between two polymers, an undesirable conjugate configuration results which may influence the splitting effect of the filament. The shape of the
fine fibre after splitting is a geometrical triangle that makes the split micro-fibre suitable for wiping application.

**Sea and island conjugate spinning process**

Polymer modification technology is used to create an alkali soluble type of copolymer. Then, the copolymer is spun with a regular polymer. The resulting fibre cross-section shows multi-islands distributed in a sea component. The sea component is then removed by the alkali soluble effect during the fabrication and dyeing process. Finally, the independent island component becomes a very fine fibre as shown in Fig. 3.19. This process is called the sea and island conjugate micro-fibres.41

The common polymer used for sea and island conjugate fibre is polyester, the denier per fibre after alkali treatment is around 0.02–0.06 denier. If a spunbonded nonwoven process is utilized, fineness may reach $9 \times 10^{-5}$
denier and the diameter of fibre is 0.1 micron. The end product from sea and island micro-fibre process could be high quality artificial leather which is used for apparel and other textiles industries. Figure 3.20 displays the scheme of spin pack for sea and island conjugate fibre (37 islands in the sea).

The appropriate conjugate ratio of alkali soluble component is 15–30%, which will be removed out during dyeing process. The flowing behaviour of this easily soluble polymer should ensure that the island component will be enveloped completely to prevent the stickiness of islands from influencing the evenness of fine fibre separation. Easily soluble polyester is developed by introducing a polymer, such as polyethylene glycol (PEG) and other monomers, to reduce resistance for the alkali solution. The alkali soluble effect of easily soluble polyester may reach 100 times that of regular polyester.
Electro-spinning process

Electro-spinning is the earliest method for the formation of nanoscale fibres.\textsuperscript{42,43} The scheme of the electro-spinning device is shown in Fig. 3.21. Polymer solution or melt is delivered by a capillary injection tube. A 30–50 kV high voltage is supplied by a power unit so that the polymer is dragged by the electric charge forming the fibre profile. Fibre from the electro-spinning process is non-oriented, so that it is hard to produce continuous filament. Thus, the fibre is formed as a nonwoven structure with diameter of fibres around $10^0$–$10^1$ nm. The tenacity of electro-spinning fibre is relatively low, which affects its application possibility.

The most significant property of polymer used for electro-spinning is viscosity and electron conductivity. The former is related to spinnability, while the latter is related to the extent of polymer dragged by the electric charge. The characteristic of electro-spinning is ease of operation for manufacturing micro-fibres with a high specific surface area of textile structure. The end product is good for industrial filtration. In addition, electro-spinning technology may have the potential for the development of tissue engineering material and artificial blood vessels for biomedical applications.

3.7 Acknowledgements

I would like to thank Dr Kuen-Shan Hwang from Acelon Chemicals and Fiber Corporation in Taiwan, Ta-Chung An from Taiwan Textile Research
Institute, Chun Lin at the Department of Materials Science and Engineering, and Abbas Al-Refaie at the Department of Industrial Engineering and Systems Management in Feng Chia University, Taiwan, for their support.

3.8 References

4.1 Introduction

Polyamides are macromolecules which contain recurring amide groups as integral parts of the polymer backbone. And nylons are polyamides with structural units derived predominantly from aliphatic monomers. Although many reactions are known that are suitable for polyamide formation, commercially important nylons have been obtained by either of two basic approaches (polycondensation and ring opening polymerization) as represented by the following general equations:

\[ \text{H}_2\text{N}\text{--(CH}_2\text{)}_x\text{--NH}_2 + \text{HOOC--(CH}_2\text{)}_y\text{--COOH} \rightarrow \text{--(HN--(CH}_2\text{)}_x\text{--NHCO--(CH}_2\text{)}_y\text{--CO--) + H}_2\text{O} \]  

Equation (4.1) refers to the synthesis of AABB-type nylons through polycondensation of bifunctional monomers utilizing stoichiometric pairs of dicarboxylic acids and diamines,

\[ \text{H}_2\text{N--(CH}_2\text{)}_z\text{--COOH} \rightarrow \text{--(HN--(CH}_2\text{)}_z\text{--CO--) + H}_2\text{O} \]  

Equations (4.2) and (4.3) pertain to the synthesis of AB-type nylons entailing respectively the polycondensation of aminoacids and the ring-opening polymerization of lactams.

The polycondensation Reactions (4.2) and (4.3) proceed by a mechanism that is characterized by what is generally known as carbonyl addition-elimination reactions which may be catalyzed or uncatalyzed.
Assuming equivalence of all the amide groups formed and independence of the functional groups from the molecular chain length, the polymerization reaches equilibrium. This may be represented by the equation

\[ -\text{NH}_2 + \text{HOOC}^- \leftrightarrow -\text{NHCO}^- + \text{H}_2\text{O} \quad [4.4] \]

The equilibrium constant \( K \) for the above reaction is defined as:

\[ K = \frac{[\text{NHCO}][\text{H}_2\text{O}]}{[\text{COOH}][\text{NH}_2]} \quad [4.5] \]

where \( B \) is the temperature independent part of equilibrium constant and \( \Delta H_a \) is the enthalpy change of the polymerization reaction. The reaction is exothermic and the \( \Delta H_a \) values in the range of \(-25\) to \(-29\) kJ mol\(^{-1}\) are reported.\(^1\),\(^2\) Therefore, a decrease in temperature favors a polymer of higher molecular weight if water concentration is constant. The molecular weight of the resulting polyamides will, thus, always remain finite and be affected considerably by any stoichiometric imbalance of the end-groups. Non-equivalence of the concentration of the bifunctional reactants and addition of monofunctional species (such as acetic acid) are practiced for molecular weight control, also known as stabilizers.

Two commercially important textile grade fibres belonging to this chemical class are nylon 6 (AB type) and nylon 6,6 (AABB type). The design of an optimal process and reactor system is possible only when it is based upon a thorough understanding of the mechanism and kinetics of the polymerization process. Therefore, a considerable amount of research has been directed to investigate the polymerization mechanisms and to determine the kinetic and thermodynamic parameters of the polyamide-forming reactions.

### 4.2 Nylon 6,6

Nylon 6,6 is prepared from polycondensation of hexamethylene diamine (HMD) and adipic acid. The repeat unit of the polymer is \(-\text{NH} - (\text{CH}_2)_6 \text{NH CO(CH}_2)_4\text{CO}^-\). The diamine, which melts at 40.87°C, is normally used in the form of a concentrated aqueous solution. The dibasic acid is used in its pure solid form (m.p. = 152.1°C). In the preparation of nylon 6,6 polymer, the first step is preparation of salt from precisely stoichiometric quantities of the intermediates at room temperature. The salt can be prepared by mixing the alcoholic solutions of the two components, the pure salt precipitates and is dissolved in water to give a solution. Alternatively, the salt can be prepared by mixing a dispersion of the diacid in water with a solution of the diamine. The aqueous salt solution is prepared so that it has a concentration of about 50–60%. The stoichiometric equivalence is
determined by measurement of the pH of a 9% salt solution prepared by dilution of main salt batch. The pH at equivalence is about 7.6 for this concentration and a shift of pH of only 0.1 results in a change in end group balance of about four equiv. $10^{-6}$ g. This small change in pH can result in unsuitable changes in the concentration of amine end groups in the polymer and fibre for those end uses where a high degree of uniformity of dyeing with acid dyes is required:

$$\begin{align*}
H_2N-(CH_2)_6-NH_2 & \quad H_3N^+-(CH_2)_6-N^+H_3 & \quad H[HN-(CH_2)_6-NHOC-(CH_2)_4-CO]_n-OH \\
\rightarrow & \quad \rightarrow & \quad \rightarrow \\
HOOC-(CH_2)_4-COOH & \quad -OOC-(CH_2)_4-COO- & \quad H_2O \\
AH Salt & \quad Nylon 6,6
\end{align*} \quad [4.6]$$

For preparation of nylon 6,6 polymer, the salt solution is subjected to evaporation at boil, possibly at elevated pressures, until concentrations $\geq 60\%$ are achieved. The concentrated salt solution together with a small amount (0.5%) of a molecular weight stabilizer is then heated in a reactor under a blanket of nitrogen so that temperature increases gradually and pressure reaches, typically, 1.73 MPa (250 psi). As water evaporates and temperature increases from about 212˚C to 275˚C, the molecular weight of the polymer reaches about 4000. Further reaction is achieved by a gradual decrease in pressure to atmospheric and then holding the polymer under these conditions for about one hour. At this point the polymer is not quite equilibrated but molecular weights are in the range of 12000 to 17000. The process is devised to remove all the liquid water present in the salt solution as well as almost all the potential water of reaction present in the form of carboxyl (–COOH) and amine (–NH$_2$) end groups with only minimal loss of the diamine (HMD) whose atmospheric pressure boiling point is 200˚C. The molecular weight of the polymer is limited by the polymer-water-steam (vapour-liquid) equilibrium at atmospheric pressure. High molecular weight nylon for industrial applications is obtained by conducting the final stages of melt polymerization under reduced pressure or by addition of chain coupling agents. The finished polymer is then extruded in the form of a ribbon or strand, quenched with water and cut to form chips. After drying, the chips are sent for subsequent spinning operation. Alternatively the polymer may be sent directly to a spinning machine without prior solidification. In nylon 6,6 the polymerization reaction goes almost to completion. The formation of cyclic oligomers is less probable than for nylon 6 since the smallest possible ring is large, with 14 members for nylon 6,6, and the end groups are, consequently, less likely to approach
each other and react. Concentrations as low as 1–2% have been reported; this level is acceptable in fibres and therefore is not removed.\textsuperscript{3}

This is represented by equation (3.29). The equilibrium constant of polyamidation reaction (equation 4.4) is shown in equations 4.5 and 4.7, where $A$ is the concentration of amide groups (–CONH–) and $P$ is the product of functional end groups (–COOH $\times$ –NH$_2$). For typically high extents of reaction and molecular weights, the value of the equilibrium constant ($K$) at 280°C is about $300 \pm 50$:\n
$$K = \frac{A[H_2O]}{P}$$  \hspace{1cm} [4.7]

At high conversions the amide group concentration is almost constant as molecular weight varies, and the water concentration in the melt at a given temperature depends only on the water vapour pressure; the equilibrium value of $P$ is proportional to water pressure. Deviation from this first power relationship is observed at high steam pressures and low molecular weights. The polymerization reaction follows second order kinetics at conversions up to about 90% and is not accelerated by catalysts.\textsuperscript{5–6} However, at the higher conversions where molecular weights become of practical interest, the reaction becomes third order and it is catalyzed by –COOH end groups and other catalysts like hypophosphite salts or phosphonic acids.\textsuperscript{7} The kinetic and equilibrium model for nylon 6,6 polymerization has been reported by Steppan and other researchers.\textsuperscript{6} In addition to –NH$_2$ and –COOH end groups, the polymer may contain stabilizer or non-functional end groups which are introduced deliberately or produced by degradation.

### 4.3 Nylon 6

Nylon 6 can be considered to be the condensation polymer of ε-aminocaproic acid with a repeat unit of –NH(CH$_2$)$_5$CO–. In fact, this polyamide is almost always prepared from ε-caprolactam in a process which is essentially an additional polymerization. The advantages of this monomer are low cost and relative ease of purification compared with the amino acid. The pure monomer, which melts at about 69°C, does not polymerize when heated at elevated temperatures in the dry state. The key discovery (11) that in the presence of both amine and carboxyl groups ring opening occurs readily, made possible the commercial processes for the polymerization of caprolactam. The simplest method is to carry out the polymerization\textsuperscript{9} in the presence of water which hydrolyzes some of the lactam to form –COOH and –NH$_2$ groups which then catalyze the addition polymerization reaction. This can be accomplished at atmospheric pressure, but the times required to achieve equilibrium monomer content can
be greatly reduced by operation at elevated pressures and increased water concentrations.

The kinetic scheme for synthesis of nylon 6 includes following three main equilibrium reactions:

(a) Ring opening

\[
\text{\(\varepsilon\)-Caprolactam (CL) \quad \text{Amino caproic acid (P₁)}\)}
\]

(b) Polyaddition

\[
\text{P}_n + \text{CL} \rightleftharpoons \text{P}_{n+1} \quad \text{[4.9]}
\]

(c) Polycondensation

\[
\text{P}_n + \text{P}_m \rightleftharpoons \text{P}_{n+m} + \text{W} \quad \text{[4.10]}
\]

where CL is caprolactam, W is water, Pₙ is polymer chain containing n repeat units. In addition, the formation of higher cyclic oligomers is an important side reaction. At equilibrium the presence of a significant concentration of the cyclic monomer along with lesser amounts of cyclic dimer and higher oligomers is known. For example, the equilibrium concentration of caprolactam at 250°C is about 7.8% and of the dimer is 1.13%.

The ring opening reaction of caprolactam is endothermic and uses water to produce aminocaproic acid which acts as a monomer for other reactions. Therefore, the rate of this reaction depends upon the concentration of water in the mixture and temperature. Without an adequate amount of ring opening, rates of reaction are very low. In the polyaddition reaction, the monomer (CL) adds on the growing polymer chain and the equilibrium of this reaction alone decides the conversion of the polymerization reaction. The polycondensation reaction is responsible for the increase in molecular weight and produces water as a condensate product. The equilibrium of the polycondensation reaction decides the molecular weight of the resultant polymer. Both the polyaddition and condensation reactions are exothermic in nature.

As explained earlier the condensation polymers require to be stabilized for molecular weight to avoid post-polymerization during processing. As shown below, the incorporation of carboxylic acid-containing compounds can result in a polymer capped at one end due to reaction with stabilizer (monocarboxylic acids):
\[ P_n + A_x \rightleftharpoons P_{nx} + W \]  \[4.11\]

where \( A_x \) and \( P_{nx} \) represent a monocarboxylic acid stabilizer and the stabilized polymer, respectively.

Kinetics and mechanistic aspects of the polymerization have been reviewed in detail\(^2,10–11\) and the rate equations and kinetic and thermodynamic constants have been used successfully for simulation and control of the hydrolytic polymerization process.

Molecular weights of nylon 6 are generally in the same range as those of nylon 6,6. For nylon 6, a commonly used viscosity measurement for molecular weight involves use of a solution of 1 g of polymer in 100 cm\(^3\) of 96% H\(_2\)SO\(_4\). A value of 2.7 has been quoted\(^{12}\) for polymer with \( M_n = 20000 \) and the value of \( \alpha \) in the Mark–Houwink equation is 0.7. For melt viscosity, \( \alpha \) was 3.5 and its low shear rate value was about 140 Pa s (1400 poise) at 280\(^\circ\)C. For nylon 6 prepared as described above, the molecular weight distribution of the final polymer is the same as for other typical condensation polymers with \( M_w/M_n = 1 + p \) or about two.

Factors influencing the polymerization and end product are temperature, water concentration and stabilizer-type and content.

### 4.4 Effect of temperature

As a consequence of the exothermic nature of the addition and condensation reactions, lower temperature is favorable for obtaining higher equilibrium values for both monomer conversion and degree of polymerization.

### 4.5 Effect of water concentration

From the mechanism and corresponding rate equation, it is readily seen that for a given temperature, the concentration of water affects the ring-opening and polycondensation reactions, and therefore, is the principal process parameter affecting both rate and attainable degree of polymerization. An increase in water concentration causes an increase in the ring opening reaction, creating more chains or end groups. The higher concentration of carboxylic acid groups is responsible for higher rates of reaction due their catalytic behavior:

\[
\text{Rate constant } k = k_i^0 + k_i^c \text{[concentration of end groups]} \]  \[4.12\]

It must be noted that water has no bearing on the equilibrium conversion because conversion is controlled by polyaddition. With increase in water concentration the molecular weight goes down, because water is a by-product; equilibrium may be shifted to higher molecular weight only on removal of water.
4.6 Effect of stabilizer type and amount

Stabilizers block the functional groups; therefore excess of one functional group may result in stabilizing molecular weight towards the end. As mentioned above, the carboxylic functional groups catalyse the hydrolytic polymerization of nylon 6 (Equation 4.12). However, since monoacid or diacid stabilizers may provide additional COOH groups, rate constants for all polymerization reactions increase in the presence of stabilizer. In recent years diacid stabilizers have gained importance in nylon 6 polymerization because such monomers may be modified to contain sulphonic acid groups, which may be used as anionic dyesite for basic dyeablenylons. The diacid stabilizers are expected to provide higher level of catalytic activity in comparison to monoacid stabilizers. This favorably affects the reaction kinetics and reduces the cost of polymerization.

The kinetics of polymerization using both monoacid and diacid stabilizers have been studied by Agrawal and coworkers. Using reaction kinetic calculations it was shown that in a closed isothermal batch reactor, the polymerization reactions proceed at a much higher rate in the diacid catalyzed system due to the presence of the two \[\text{COOH}\] groups in its chemical structure. On the other hand, monoacid has only one \[\text{COOH}\] group in its chemical structure, and therefore, shows lower catalytic activity. It is well known that the carboxylic (–COOH) groups catalyze all the kinetics reactions involved in the polymerization of caprolactam. Although both monoacid and diacid work as a stabilizer in controlling the molecular weight of the polymer at the end of the polymerization, the higher rates of conversion and molecular weight build up are observed in the case of diacid stabilized system.

Figures 4.1–4.3, respectively, show the comparison of the two systems polymerized at 265°C for build up of conversion, carboxylic groups, and amino groups with respect to the reaction time. It can be observed that from the beginning of the reaction, the concentration of the carboxylic end groups in the case of a diacid stabilized system is higher than in a monoacid stabilized system. The difference is maintained throughout the reaction and results in higher catalytic activity for all polymerization reactions. The concentration of amino end groups rises faster resulting in a peak occurring earlier to that in monoacid stabilized system. The concentration of amino end groups in case of monoacid stabilized system reaches maximum at about 1.0 h while in the case of a diacid stabilized system it is about 0.5 h. Table 4.1 shows that for both diacid and monoacid stabilizers, the conversion reached similar values, which confirmed that the conversion is independent of the type or concentration of stabilizer used.
4.7 Reactor design

Nylon 6 polymerization may be carried out in a semi-batch reactor or a continuous tubular reactor (VK tube).

4.7.1 Semi-batch reactor

From the understanding of the factors affecting the kinetics of the polymerization reaction in nylon 6, it may be easy to infer that it is desirable...
to alter the condition of the polymerization during the initial and the final phases of the reaction. Initially, the water is kept high by applying high pressure in the reactor to facilitate ring opening reaction. Also the temperature of the reaction is kept high to allow rapid conversion of the monomer and build-up of the molecular weight. At the later stages, the reactor is slowly brought under low pressure and to lower temperature to favour high molecular weight and high conversion.

This scheme of polymerization may also be carried out in a semi-continuous manner by using a sequence of two semi-batch reactors. The first reactor should have a higher temperature and higher water concentra-
tion so as to achieve a rapid reaction rate and the second should have a low temperature and low water concentration in order to drive the polycondensation and polyaddition reactions in the forward direction to obtain high molecular weights and conversions.

### 4.7.2 VK tube

One of the popular reactors used in industries for the hydrolytic polymerization of ε-caprolactam is the VK (Vereinfacht Kontinuierliches Rohr) column reactor. VK column reactors are specially designed vertical tube reactors where internal gratings are placed in order to achieve uniform heating and near flat velocity profile of reaction mass.

The industrial VK tube reactor consists of several different zones, where the temperature is independently measured and controlled. A schematic representation of the reactor with five zones is given in Figure 4.4(A). Feed consisting of molten caprolactam, water and mono- or di-functional acid is fed under the nitrogen cover at the top of the column. The purged nitrogen is made to escape through a water seal located at the top of the reactor. The reaction mixture fills the entire tube and is heated in zone I as it passes through a heat exchanger I to a temperature of about 240°C. Thereafter (zone III upper and lower) the melt is heated by the heating jackets and the ongoing exothermic reactions. The temperature of the reaction mass gradually increases from the top to the middle zone till a highest temperature point (Th) of about 260–270°C is reached. Thereafter, the temperature is decreased to 240–250°C using another heat exchanger and stabilized towards the bottom of the column using the jacket temperature. The products, polymer and unreacted monomer, are removed at the bottom of the reactor.

As stated above, ring-opening and polycondensation reactions are very sensitive to the water content in the reaction mass at any point. Use of water also affects the concentration of carboxylic end groups, which helps in catalyzing all the polymerization reactions. Therefore, water concentration profile in reaction mass along the axis of VK tube reactor significantly affects the end product properties.

In a VK tube reactor, large quantities of water are added to initiate the ring-opening reaction. The high temperature, at the top portion of the VK tube, readily evaporates the volatile components, water and caprolactam, until equilibrium is attained between the vapor and liquid phases. Evaporated caprolactam is condensed and returned back to the VK tube. Most of the added water is evaporated and removed as condensate within a short time after addition. As the reaction mass moves down the reactor towards the highest temperature point (Th), more and more water is evaporated, and the vapor rises to the top surface of the reactor. This transfer
of water vapor helps in maintaining both the liquid and vapor phases in near equilibrium state at all places in the upper portion of the reactor. This gradual evaporation of water due to rising temperature results in a 'lowest water content (LWC)' point.

Since the water can vaporize and escape by rising to the surface of the melt, the water content at any point in the reactor depends on the combined action of temperature and pressure till the lowest concentration of water is reached. Higher temperature tends to reduce water content whereas higher pressure increases it. This counteraction of the two parameters results in a point in the VK tube that is the 'lowest water content (LWC) point'.
When the reaction mixture crosses the highest temperature point ($T_h$),
the lower temperature and higher hydrostatic pressure in the bottom
portion of the reactor suppresses the vaporization of water, and the remain-
ing portion of VK tube reactor is practically a non-vaporizing tubular
reactor. Therefore, a VK column reactor can be considered as a reactor
consisting of three main zones: at the top is a turbulent zone, which may
be considered as a two-phase (gas-liquid) continuous stirred tank reactor
(CSTR) type zone, a middle zone as a vaporizing plug flow tubular zone,
and a bottom zone as a non-vaporizing plug flow tubular zone. The three
zones are shown in Figure 4.4(B).

The VLE equation suggested by Fukumoto\textsuperscript{17} for the end of polymeriza-
tion under vacuum conditions has been found to be suitable for most of
the vaporizing zone based on the comparison between the predicted and
experimental values:

$$[W] = 10^{-2} P_r \exp \left( \frac{8220}{T} - 24.36 \right)$$ \hspace{1cm} [4.13]

The prediction of polymer properties is improved by modifying the
constants in the equation to account for the pressures higher than the
atmospheric pressure condition inside the VK tube:

$$[W] = 10^{-2} P_r \exp \left( \frac{8220}{T} - 22.24 \right)$$ \hspace{1cm} [4.14]

where $P_r$ is the total pressure in Pascal defined by

$$P_r = P_A + 9.81 (\rho_w H_w + \rho_m H_p)$$ \hspace{1cm} [4.15]

where $P_A$ = atmospheric pressure $= 1.013 \times 10^5$ Pa.

Equation 4.15 can be used for calculating concentration of water at every
point in the vaporizing zone of the reactor. From the ‘lowest water content
(LWC)’ point downward (i.e., non-vaporizing zone), the summation of
water in the reaction mixture, i.e. $[W]$, and the water which is part of the
polymer i.e. $[P]$, can be considered to be constant. The concentration of
water $[W]$ at any point in non-vaporizing zone is then estimated using a
mass balance equation as follows:

$$[W] + [P] = [W_c] + [P_c]$$ \hspace{1cm} [4.16]

where $[W_c]$ and $[P_c]$ are the concentrations of water and polymer at
the ‘lowest water content (LWC)’ point, respectively; $[P]$ is the concen-
tration of polymer at the point of consideration in the non-vaporization
zone.

Figure 4.5 shows the temperature and pressure variation along the axis
of a typical VK tube reactor. The pressure at any point inside the reactor
is a summation of atmospheric pressure acting on the top of the reactor, head pressure due to the water seal length at the top of the reactor, and the hydrostatic pressure caused by the weight of the polymer mixture on the top of the concerned point. The pressure on the reaction mixture changes from $1.023 \times 10^5$ Pa at the top of the reactor to $2.082 \times 10^5$ Pa at the bottom of the VK reactor of about 10–11 m height. Besides the above, the flow of viscous polymer melt through the VK tube may also lead to pressure losses which need to be accounted for if large. However, such pressure losses are expected to be small and therefore can be neglected. The pressure variation is generally non-linear with the reaction time because the effective cross-sectional area of the reactor is different in different reaction zones.

The counter-flow direction of the rising water vapor in the vaporizing zones can cause some agitation of the reaction mass. Due to this reason, the earlier models of VK tube\textsuperscript{11,16,17} assumed the vaporizing zones to behave like a CSTR and modeled the VK column reactor as CSTR (or a series of CSTRs) followed by a plug flow reactor. However, modeling the entire vaporizing zone of VK tube as a combination of CSTRs may be incorrect because significant agitation exists only in the top zone of the reactor. Figure 4.6 shows a profile of the water concentration inside a typical VK tube. It can be observed from the figure that the most of the water evaporates soon after it is added at the top of the reactor. This occurs within 0.5 hours of the start of the reaction. Therefore only this first 0.5 hours of the reaction may be considered as a CSTR.
In a VK tube reactor pressure plays a crucial role. The increasing pressure inside the VK tube along with temperature define the concentration of the water available for the reaction. The pressure inside the VK tube may be altered by applying a static pressure at the top of the reactor. Figure 4.7 depicts the effect of head pressure (which is the pressure created by the water seal at the top of the VK tube) on the number average molecular weight of the polymer. As an example, increase in water seal length from

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**Effect of pressure in a VK tube**

In a VK tube reactor pressure plays a crucial role. The increasing pressure inside the VK tube along with temperature define the concentration of the water available for the reaction. The pressure inside the VK tube may be altered by applying a static pressure at the top of the reactor. Figure 4.7 depicts the effect of head pressure (which is the pressure created by the water seal at the top of the VK tube) on the number average molecular weight of the polymer. As an example, increase in water seal length from...
0.1 m to 5 m, results in significant improvement in the rate of reactions as more water may tend to stay inside the reactor. However, due to larger water content at the equilibrium, number average molecular weight decreases by about 10.5% and amino end groups increase by about 15.5%.

**Effect of temperature profile in the VK tube**

The temperature profile in the VK tube reactor is another important parameter in optimizing reactor performance. It is important that the temperature of the reaction mass is first increased slowly to the highest temperature point and then decreased to the final desirable temperature of equilibrium. This profile helps in first retaining the water for proper initiation of the reactions and later allows removal of water to the desirable extent at the highest temperature point. Water content at the highest temperature point ultimately decides the water content at the end of the reaction, and thereby, the properties of the polymer.

The temperature at the ‘highest temperature point (T_h)’, therefore, plays an important role in deciding the polymer properties. When the highest temperature is allowed to increase, the water content at the lowest water point decreases. This decrease in [W] in the non-vaporizing zone, increases the number average molecular weight as depicted in Figure 4.8. The calculations show that with a nominal change of the highest temperature (T_h) by 10 K, the molecular weight changes significantly by about 11.5%.

![Figure 4.8 Effect of the highest temperature (T_h) on the number average molecular weight.](image-url)
**Effect of stabilizers**

In a VK tube reactor, the use of diacid stabilizer over monoacid stabilizers also brings in substantial change in the kinetics of the reaction and the overall reaction time. Due to higher rates of reactions, diacid stabilized system achieves equilibrium faster than monoacid stabilized system and results in shorter polymerization time by 20–25%.15,16

**Effect of internal design of the VK tube**

It is important to control both the pressure and temperature profiles in a VK tube reactor in order to achieve desirable reaction time. It is desirable to have high pressure at relatively high temperature in the vaporizing zone. This would allow polymerization to reach equilibrium at a faster rate. Eventually, added water may be removed to the desired extent by increasing the temperature at the highest temperature (T_h) point before cooling the reaction to a lower temperature in the non-vaporizing zone. However, in a conventional VK tube, the possibility of increasing T_h is limited by the low heat-removal capacity of the reactor. Uncontrolled heat generation at the center of the reactor due to fast exothermic reactions at high temperature may result in gelling or degradation of the polymer. This limitation may be overcome by providing an efficient cooling device such as an additional heat exchanger just above the highest temperature point. This would allow controlled polymerization at the desirable temperature profile.

Other variations in the designs of VK tube involve a gas purging device at the end of the vaporizing zone to control the concentration of water at the lowest water content point. For industrial nylon 6, the molecular weights may be increased by adding a vaporizer.

**Conversion of monomers in VK tube**

Although it is not necessary to reach an absolute equilibrium in nylon 6 polymerization, near equilibrium conditions are desirable for a stable system with dependable results. The extent of a reaction may be determined by studying ‘Ψ_i factor’, which is defined for a reaction ‘i’ by the following expression:

\[
Ψ_i \text{ factor} = \frac{(R_{if} - R_{ib})}{R_{if}} \quad i = 1, 2, 3
\]

where \( R_{if} \) and \( R_{ib} \) are forward and backward rates of reaction for a given reaction ‘i’. A reaction reaches equilibrium when the ‘Ψ_i factor’ becomes zero, in other words, the forward reaction rate becomes equal to the backward reaction rate. The ‘Ψ_i factor’ versus reaction time for all the
major reactions – ring-opening, polycondensation and polyaddition are plotted in Figure 4.9 for an experimental case. It is interesting to note that all reactions show a similar trend. Initially at the start of polymerization, all reactions show $\Psi_i = 1$ as there is no possibility of backward reaction. The ‘$\Psi_i$ factor’ decreases with time for all reactions, except for the ring-opening reaction, which shows a sudden increase in $\Psi$ at about 0.5–1 h. This increase in $\Psi_i$ may be due to a lower rate of backward reaction as aminocaproic acid is utilized by the other two reactions. All reactions tend to reach equilibrium together at about 7–8 h. However, $\Psi$ values for polycondensation and polyaddition reactions increase when the reaction equilibriums are disturbed favorably by lowering the reaction temperature in the non-vaporizing zone. However, ring-opening reaction continues its downward trend at this zone since it is an endothermic reaction.

Net rates for all reactions start to increase almost simultaneously. The maximum value for the net rate of reaction for ring-opening occurs at around 2.3 h followed by polyaddition at around 2.8 h and finally polycondensation at 3.4 h. The peak values for all the three reactions occur at reaction times that are close to each other and are within the initial 30% of the overall reaction time. The rate of reaction for polyaddition is higher than those of the other two by about one–two orders of magnitude. From Figure 4.10, it can be seen that curve of polyaddition overlaps the curves
of ring-opening and polycondensation from the beginning to the end of the reaction.

The concentration of functional end groups also changes with reaction time (Figures 4.11 and 4.12). The end group concentration is maximum at around 4 h due to the higher rate of ring-opening reaction than polycondensation. Thereafter, polycondensation rate predominates (Figure 4.10) over the former and results in the decrease of end group concentration. All this time polyaddition is the fastest reaction of all, helping to improve the molecular weight of the polymer at a slow and steady pace. However, it must be kept in mind that increase in molecular weight is rather poor with polycondensation reaction. The final molecular weight is achieved only because of polycondensation reaction.

4.8 Synthesis of modified polyamides (nylon 6)

Nylon 6 may be modified during the polymerization with comonomers or stabilizers to introduce new functional groups or chain end groups and thus
4.11 Calculated amino end group \([\text{NH}_2]\) concentration in a typical VK tube: (A) considering the effect of increasing hydrostatic pressure inside the reactor; and (B) assuming constant atmospheric pressure throughout the reactor.

4.12 Calculated carboxylic acid end group \([\text{COOH}]\) concentration in a typical VK tube: (A) considering the effect of increasing hydrostatic pressure inside the reactor; and (B) assuming constant atmospheric pressure throughout the reactor.
change the reactivity and chemical properties such as dyeability. In addition, the polyamide fibres can also be subjected to a variety of treatments including chemical, grafting, radiation, and plasma which alter the surface characteristics of these polymers.

Various nylon 6 nanocomposite polymers have been synthesized by in situ polymerization ε-caprolactam in the presence of nanomaterials.\textsuperscript{18–21} Nylon 6/nano-TiO\textsubscript{2} and nylon 6/nano-Al\textsubscript{2}O\textsubscript{3} composites have excellent photooxidative degradation resistance.\textsuperscript{18,19} In the presence of pristine and carboxylated multi-walled carbon nanotubes (MWNT and MWNTCOOH) both the storage modulus (\(E'\)) and glass transition temp. (\(T_g\)) of the PA6/CNTs composites increase significantly.\textsuperscript{20,21}

4.9 Modification at polymerization stage

Nylon 6 may be modified during polymerization by introducing small amount of a comonomer containing desirable functional side groups.

Since –NH\textsubscript{2} and –COOH are the functional groups involved in the polymerization reactions, a comonomer may be added with the following general chemical formula for synthesizing a copolymer with nylon 6:

\[
\begin{align*}
\text{H}_2\text{N} & - \text{R} - \text{C} - \text{OH} \\
& \quad \text{R}'
\end{align*}
\]

Alternately, two monomers may be added in equal molar ratio as given below:

\[
\begin{align*}
\text{H}_2\text{N} & - \text{R} - \text{NH}_2 \\
& \quad \text{R}' \\
+ & \quad \text{HO} - \text{C} - \text{R}'' - \text{C} - \text{OH} \\
& \quad \text{R}''
\end{align*}
\]

Apart from using comonomers to introduce new functional groups, stabilizers such as monoacids, monoamines, diacids or diamines, may also be used for introducing desired functionality to nylon 6. These stabilizers modify the end groups of the polymer chains and thus change their reactivity and chemical properties such as dyeability:

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Another approach that is used for modifying nylon 6 is by transamidation reaction. In this method, the tendency of polyamides to break and reform their amide linkages is utilized. After breakage, the amide linkage may reform with a free amine of any available polyamide segment. A typical representation of such reaction is:

\[ \text{Monoamine} \rightarrow \text{Diamine} \]

\[ \text{Monoacid} \rightarrow \text{Diacid} \]

Using the above reaction mechanism, a new group $R''$ is readily introduced in nylon 6 by mixing another suitable polyamide containing $R''$ at the polymerization temperature. Since the above exchange occurs randomly at any amide linkage, initially after lapse of short reaction time, the two polyamides react to give a block copolymer with segments of each polyamide. If the transamidation is allowed to continue for a longer time then these segments further undergo amide exchange and eventually result in a random copolymer as shown in Figure 4.13. Therefore, care must be taken to arrest the reaction at an appropriate time so that only the desired amount of segmental exchange takes place. A completely random
4.10 Dyeability of nylon 6

ε-Caprolactam, if allowed to polymerize without the use of a stabilizer, results in a polymer with equal numbers of –NH₂ and –COOH groups. For a polymer of about 16000 g/mol molecular weight, end group concentration is about 62 eq/ton for each type of the two end groups. Addition of a small quantity of stabilizer such as acetic acid blocks some number of NH₂ groups and helps in producing nylon 6 with normal dyeability, where an NH₂ end group concentration of about 35–45 eq/ton is targeted. Since NH₂ groups are the reactive functional sites for attracting acid dyes, any variation in amino end-group concentration changes the dyeing characteristics of the resulting fibres. If more NH₂ groups are blocked by adding a little more monoacid or a diacid such as sebasic acid, then dyeability of nylon with acid dyes may decrease substantially. A concentration of 10–15 eq/ton of NH₂ groups in the final polymer results in light dyeable nylon 6.

Similarly, if the concentration of NH₂ groups is increased beyond 40 eq/ton to the values of 80–90 eq/ton, deep dyeable nylon 6 is achieved. This increase in NH₂ concentration may be obtained by adding required quantity of diamines in place of mono or diacid as stabilizers.
Diamines stabilize the molecular weight by reacting at the –COOH end group, and altering the 1:1 ratio of amino and carboxylic end groups. However, in doing so they convert the –COOH end group to an additional NH₂ end group. This results in a much higher concentration of NH₂ end group than possible with unstabilized/unmodified nylon 6:

\[
\text{H}_2\text{N} \equiv \text{COOH} + \text{H}_2\text{N} \equiv \text{NH}_2 \rightleftharpoons \text{H}_2\text{N} \equiv \text{C} - \text{NH} - \text{NH}_2
\]

Nylon 6  Diamine stabilizer  Modified nylon 6

The dyeability of nylon 6 may be further enhanced and grades such as ultra deep (with NH₂ = 95–100 eq/ton) and super ultra deep (with NH₂ = 115–130 eq/ton) may be produced using a greater quantity of diamines to block additional –COOH end groups mentioned above.

The addition of a much higher quantity of stabilizers either mono/di acids or diamines for obtaining required dyeability characteristics places
stringent regulations on polymerization process. Obviously, the molecular weight of modified polymer may become substantially lower if the condensation reaction is not pushed forward by removing additional quantity of free water. Alternately, dyeability of the nylon 6 may be modified by adding a second polyamide flake of the same polymer type but adding a different amine end level to the extruder with the first polyamide flakes. This is followed by mixing and melting the two flakes under specific conditions to produce a blend with the required concentration of amino end group.

4.11 Cationic dyeable nylon 6

Nylon 6 is principally dyed using only the acid dyes as amino groups are highly active dye sites. The other functional group, carboxylic end group, forms only a weak acid group and therefore cannot attract basic or cationic dyes in normal nylon 6. For achieving cationic dyeability, stronger functional groups such as sulphonate acid groups are required to be introduced during polymerization. Use of modified stabilizer containing cationic dye sites has been suggested. Disodium 3,5-disulphobenzoic acid with following structure may be used as the stabilizer to the extent of about 1 wt% on caprolactam22;

\[
\text{HOOC} - C_6H_4(SO_3Na_2) - COOH + H_2N \rightarrow \text{nylon 6}
\]

The above molecule reacts at the amino end group and provides two sulphonate groups as shown below:

\[
\text{H}_2\text{N} - COOH + \text{HOOC} \rightarrow \text{C}_6\text{H}_4(SO_3\text{Na}_2)
\]

Cationic dyeable nylon 6 is normally less receptive towards acid dyes as amino groups are sacrificed in favour of sulphonate groups. Cationic dyeable nylon 6 may also be produced by adding 1–5% sulphonated diamino stilbene23 during the synthesis stage. It is also possible to produce a
copolymer by adding a dicarboxylic acid monomer in addition to the above
diamine mentioned. The mechanical properties of the modified nylon 6
may be improved but the improvement depends upon the concentration of
comonomers used.

4.12 Antistatic and hydrophilic nylon 6

Nylon 6 has a moderate moisture regain of about 4–4.5% at standard con-
dition. This is significantly better than many synthetic polymers such as
polyesters, polyolefins, etc., however, still it is lower than many of the
natural fibers such as cotton, silk or wool. The comfort properties of a
fabric depend on several attributes of which moisture absorbency is one of
the most important properties for applications related to innerwear and
sportswear. The moisture wicking property of fabrics, which contributes
towards comfort value under heavy perspiration conditions, has been found
to be higher for more hydrophilic fibers than hydrophobic fibers.24

Moisture regain in nylon 6 is due to the presence of hydrophilic amide
linkages. If hydrophilic groups per unit weight of the material could be
increased, then it would lead to higher moisture content at the standard
conditions. For example, nylon 4, which has a higher concentration of
amide linkages in the polymer mass, shows 9.1 wt% moisture regain. Based
on similar reasoning nylon 6 is made more hydrophilic by copolymerizing
the polymer with copolymer segments containing hetero atoms such as
ether links of poly(ethylene glycol) segments22 having the structure shown
below:

\[
H_2N-\text{CH}_2-\text{CH}_2-\text{CH}_2-O\text{-(CH}_2-\text{CH}_2-O)_n\text{CH}_2-\text{CH}_2-\text{CH}_2-NH_2
\]

Diamine functionality is used in such comonomer segments to facilitate
copolymerization with carboxylic acid end groups of nylon 6 chains. This
gives block copolymer structure. Random copolymerization using small
polyether molecules is not advisable as it may lead to formation of essen-
tially amorphous material, which is unsuitable for fiber formation.

Example of such a hydrophilic nylon 6 fiber is ‘Hydrofil’ developed by
Allied Fibers.24 This fiber is obtained from a block copolymer of nylon 6 and
polyethylene glycol with about 15% PE and rest nylon 6. At standard condi-
tions of 65% relative humidity and 70°F temperature, the fiber has moisture
regain of about 5.4% which increases to about 14.4% at 95% RH and 90°F
(32°C) temperature. The moisture absorption at higher relative humidity
compared well with that of cotton. The material is suitable for making
active wear fabrics. Alternately, a copolymer of adipic acid-caprolactam-
polyethylene glycol25 may be used as an additive in the nylon 6 matrix to
impart antistatic properties even at a relative humidity below 45%.
Hydrophilic segments such as polyether-amides may also be introduced by transamidation. The fibers produced from such composite materials have lower crystallinity and higher moisture regain. Initially, on the physical mixing, the two polyamides are in separate phases, however as time proceeds, mixers of molten polyamide undergo transamidation (amide interchange) reaction to form a block copolymer. Vivrelle polymer developed jointly by Sun Co. and SNIA fibers\(^{26}\) is an example of such a block copolymer.

### 4.13 Flame retardant nylon 6

Nylon 6 can catch fire in normal atmosphere as its Limiting Oxygen Index (LOI) is around 21. However, it has self-extinguishing properties due to its extensive shrinkage and dripping behavior during combustion. When the nylon polymer is ignited, it melts and then the molten polymer drips away from the flame. However, if the molten polymer is not able to escape fire, then it propagates fire. The nylon 6 can be rendered flame retardant by either using certain phosphorous/halogen based comonomers during polymerization or by adding flame retardant additives during melt spinning.

Bis(2-carboxy ethyl)methyl phosphine oxide (CEMPO)\(^{27}\) when used as a comonomer with hexamethylene diamine (HMDA) gives a flame retardant copolymer fiber with LOI of \(\sim 25\%\). The fibre properties are not significantly affected up to a concentration of 35 mol\% of CEMPO. The chemical structure of the additive is:

\[
\begin{align*}
O & \quad O \\
\text{HO} & \quad \text{CH}_2 \quad \text{CH}_2 \\
\text{C} & \quad \text{CH}_2 \quad \text{CH}_2 \\
\text{P} & \quad \text{CH}_2 \quad \text{C} \\
\text{OH} & \quad \text{CH}_3
\end{align*}
\]

(CEMPO)

Net phosphorus content of about 1.5 wt\% on polymer weight gives adequate protection against fire. Halogen based monomers or additives, though effective, are no longer recommended because of the emission of halogen acid during burning. Vapours of these acids may cause personal injuries.

### 4.14 Elements of melt spinning process of nyons

A fiber or a filament (a continuous form of fiber) is the fundamental unit of textile materials. It has high strength (tensile, bending, torsional, or
compression), high flexibility (i.e. low moduli), extensibility, and shows recoverability on deformation. Most of these properties are observed about one principal direction, which is known as the axis of the fiber. Since all textile structures – one to three dimensional (yarn, fabric, or braids, etc.), are built using this basic structural unit, these structures also possess such unique properties.

In order to possess such properties, the fiber or filament has a unique micro-structure (morphology) in which majority of the polymer chains are oriented in the direction of the axis of the fiber. The more oriented the polymer molecules are in this direction, the better properties the resulting fiber/filament is considered to have. For producing man-made fibers, the polymers (either natural or synthetic) must be unfolded and extended uni-directionally to extremely large dimension to get high aspect (length to diameter) ratio and high orientation. This is known as spinning.

In spinning, a small amount of polymer (say ~1 g) may be elongated to over 9000 km while the other dimension (diameter) is only in microns. This requires a precise control over the spinning process to enable such uni-directional extensions in melt form. In fiber formation, all efforts are directed in controlling the microstructure of the polymer so that properties as mentioned above are obtained with respect to the principal axis of the fiber. This is unlike other methods of polymer processing such as injection molding, compression molding, extrusion and blowing, etc., where mostly isotropic properties are desired and only a little attention is directed in making a controlled microstructure in terms of molecular orientation and their spatial arrangement.

In spinning, polymer melt or solution is extruded from a fine hole and is elongated by applying a tensile external force on the extruded portion. As the polymer melt or solution is pulled, it is cooled or precipitated, respectively, to form a solid filament. This filament is then usually subjected to post-spinning operations such as drawing, which is unidirectional stretching in a semi-solid form, and heat-setting, which is crystallization to equilibrium. Other post-spinning processes, such as texturing, simply are variations of the drawing and heat-setting processes to impart curvilinear shape to an otherwise straight filament. This gives physical bulk to the filaments. The process of fiber formation is complete only when both spinning and post-spinning operations are carried out.

When polymer is moving inside a confined geometry, it is under a shear flow such as through a pipe or spinneret hole. During this flow, the molecules of the polymer do not orient significantly and the shear flow does not contribute to formation of fiber in typical nylons. However, liquid crystalline polymers may be transformed and oriented during a shear flow through a spinneret. In a flexible chain polyamide, the spinning and structure formation takes place only after the polymer melt has been extruded.
through the spinneret. As the polymer comes out from the confined channels into the open atmosphere, i.e. extruded from the spinneret, its flow behavior changes. It is said to undergo elongational flow under tensile forces.

In an elongational flow, polymer fluid flows with a uniform velocity profile perpendicular to the direction of fluid flow. However, it may accelerate, or at other times, decelerates as it travels along its path. Most of the spinning is said to occur due to the elongational flow where fluid velocity keeps on increasing down the spinning line, resulting in unfolding of the polymer chains.

### 4.15 Structure development during melt spinning of polyamides

#### 4.15.1 Spinning at low speeds

At lower spinning speeds, stress induced crystallization is absent. Therefore, with increasing level of spinning stress, the orientation of the polymer chains increases. However, it does not have any significant effect on crystallization.

Temperature profile changes both the orientation and crystallinity of the spinning filament. Higher temperature increases the spinning length (time spent by the spinning filament in spinning line) as the filaments take longer to cool down. It tends to increase polymer chain relaxation, and therefore, lowers the orientation. At the same time, the crystallinity of the filament increases because it spends longer at a higher temperature.

The winding (spinning) speed is critical in deciding the level of spinning stress inside the spinning filament. The stress in turn regulates the velocity gradient profile of the spinning line. Velocity gradient profile is the change in velocity of the filament with spinning distance. Below a critical stress level, the velocity gradient profile is a bell shaped curve as shown in Figure 4.14. The maximum velocity gradient is obtained somewhere in the middle of the spinning line and its maximum value increases gradually with the increasing spinning speed. The orientation in the spun filament is the function of this maximum velocity gradient value, i.e. the maximum strain rate.

The temperature of the spinning line also depends on the spinning speed. At higher spinning speeds, the temperature of the spinning filament may be at a slightly lower temperature. This may be due to the enhanced cooling effect of the quench air at higher relative speed of the filament.
Throughput also has significant effect on the temperature, stress, and velocity gradient profiles. With higher throughput, the value of maximum velocity gradient decreases and it maximizes at a later distance from the spinneret. This is shown in Figure 4.15. At higher throughput, the temperature of the filaments remains at a higher temperature for a longer time.\textsuperscript{28} This is because with higher throughput (W), more heat comes into the spinning line and it takes longer for the same cooling system to cool down the fiber to its $T_g$. Therefore, not only does the overall temperature of spinning line increase, but the spinning length increases. The spinning fiber spends longer in the spinning line. The spinning length is defined as the distance traveled by the extruded fiber before it reaches a temperature below its $T_g$. Below $T_g$, the spinning is considered to be finished because
the fiber enters the glassy state and can no longer be stretched under the applied stresses.

Similarly, cooling air conditions have direct effect on temperature profile of the spinning filament, and therefore, on spinning stress. If the relative humidity of the cooling air is high, it can carry more heat due to its high heat capacity and provide better cooling to the filament. On the other hand, if the spinning polymer is sensitive to moisture such as polyamides, the moist air may act as a plasticizer and influence the $T_g$ of the polymer. This will tend to lower spinning stresses (lower orientation), but facilitate crystallization at the same time. Therefore humidity of cooling air in such systems is regulated to towards lower value. Table 4.2 summarizes the effect of spinning parameters (assuming that only one parameter is being changed at a time) on the structure of the as-spun fibers at low spinning speeds.

### 4.15.2 Spinning at high speeds

As the spinning speed is increased further to higher values, the effect of air drag and inertial forces increases substantially on the total stress value of the spin-line. Many studies have been reported in the literature on modeling of the high speed spinning process to explain the spin line dynamics.\textsuperscript{29–34} Figure 4.16 shows the effect of the spinning speed on the developed stresses of the spinning line in nylon 66.\textsuperscript{30,35}

When these stresses cross a critical stress level, which is unique for a particular polymer, the polymer starts to undergo fast deformation (high strain rate) and the stress level rises rapidly due to diameter attenuation. At the high strain rate, its apparent elongational viscosity starts to decrease rapidly due to strain thinning, which further induces even faster deformation. This is known as region of ‘super drawing’ or ‘necking’. This deformation cycle continues till polymer chains are pulled into high orientation and their degree of freedom is locked due to initiation of crystallization. This is known as stress-induced crystallization. The oriented and partly crystallized polymer chains turn into a rubber like structure and start to show strain-hardening behavior, which arrests further deformation of the structure. On strain hardening, the apparent viscosity starts to rise very sharply. This phenomenon is predicted in Figure 4.16 for nylon 66 case. At higher spinning speeds, the initial drop in the apparent viscosity is higher and faster followed by even faster rise after the stress-induced crystallization takes place. The freeze point is achieved soon after this strain-hardening phenomenon takes place. In contrast, no such changes in apparent viscosity are seen in the case of 1000 m/min of spinning speed.

The higher spinning speeds give rise to the higher necking tendency, which leads to more orientation and higher crystallinity. With higher
Table 4.2 Effect of spinning parameters on the structure of as-spun fibres

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Effect on spinning stress</th>
<th>Effect on filament temperature</th>
<th>Time spent in spinning line</th>
<th>Fibre orientation</th>
<th>Fibre crystallinity</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>↑Winding speed</td>
<td>↑</td>
<td>↓</td>
<td>↓</td>
<td>↑</td>
<td>↓</td>
<td>Higher denier</td>
</tr>
<tr>
<td>↑Throughput</td>
<td>↓</td>
<td>↑</td>
<td>↑</td>
<td>↓</td>
<td>↑</td>
<td>Slower cooling</td>
</tr>
<tr>
<td>↑Cooling air temp</td>
<td>↓</td>
<td>↑</td>
<td>↑</td>
<td>↓</td>
<td>↑</td>
<td>Faster cooling</td>
</tr>
<tr>
<td>↑Cooling air speed</td>
<td>↑</td>
<td>↓</td>
<td>↑</td>
<td>↑</td>
<td>↓</td>
<td>Faster cooling carries more heat</td>
</tr>
<tr>
<td>↑RH (hydrophobic fiber)</td>
<td>↑</td>
<td>↓</td>
<td>↓</td>
<td>↑</td>
<td>↓</td>
<td>Moisture plasticizes the polymer</td>
</tr>
<tr>
<td>↑RH (for hydrophilic fibre)</td>
<td>↓</td>
<td>↓</td>
<td>↑</td>
<td>↓</td>
<td>↑</td>
<td></td>
</tr>
</tbody>
</table>
spinning (winding) speed, the neck moves towards the spinneret and the change in velocity gradient is much steeper. The spinning is completed much sooner with the spinning path considerably reduced. Table 4.3 shows that the necking starts at a higher temperature at higher spinning speed for the three polymers studied.\textsuperscript{36} The polymer spends a very little time in the necking region. It is predicted that necking in nylon 66 occurs for only 8–10 cm at 5500–6500 m/min and the time spent by the polymer in the neck region at these speeds is only about 5 ms. The stress induced crystallization as observed above may take place only if two parameters are favorable – the induction time for crystallization (i.e. inverse of nucleation
rate) and the rate of crystallization (i.e. rate of crystal growth). For an effective stress-induced crystallization to occur at higher spinning speeds, the induction time for crystallization should reduce and rate of crystallization should increase for the spinning polymer. In other words, the overall rate of crystallization should increase.

In nylon 6, the POY speeds are nearly 4000–4200 m/min. Nylon 6 spun at lower speeds undergoes post-spinning crystallization on conditioning. Therefore, it is difficult to obtain amorphous nylon 6 at any speed due to this phenomenon. At speeds near to 4000 m/min, nylon 6 POY is stable and essentially consists of $\gamma$ crystals, which are deformable during drawing and texturing.

### 4.16 Spinning of nylon 6

In the spinning thread, in the absence of crystallinity, the polymer behaves as a single phase rubber-like network, adhering to affine deformation similar to spinning of PET. Due to a post-crystallization process that occurs after winding and conditioning, nylon 6 fibers develop a substantial level of crystallinity (50% or more) even at low wind-up speeds. The developed crystals are of $\alpha$ type. The structure of the amorphous network developed in the spinning thread is preserved in the final wound sample in spite of post-crystallization. The orientation of the crystalline phase that develops during post-crystallization is relatively high even at low wind-up speed. Apparently, post-crystallization starts from highly oriented amorphous segments, such as short network chains that become fully extended at low overall network deformation. Post-crystallization leads to a pronounced difference in birefringence determined on-line and after winding.

The change in birefringence on conditioning is indicative of the degree of instability the as-spun material shows (Figure 4.17). Upon conditioning, the fiber spun at low spinning speeds extends owing to crystallization, which throws the molecular chains in the axial direction. At a speed of 2000 m/min, the extension is the maximum while it reduces slowly to a negligible value as the spinning speed increases to about 4000–4200 m/min. At this speed, the extension due to crystallization and contraction due to residual spinning stresses are well balanced. At spinning speeds higher than 4200 m/min, the as-spun fiber shows net contraction, which increases slowly with increasing spinning speed.

Figure 4.18 shows a series of true stress–strain curves for undrawn yarns obtained over a wide range of wind-up speeds (500–6500 m/min), that have been shifted horizontally to coincide with the 500 m/min LOY curve. For yarns spun at wind-up speeds below 4000 m/min, a good fit with the mastercurve is obtained. For yarns spun at wind-up speeds above
4.17 Change in birefringence with conditioning time of nylon 6 with a mass throughput of 2.99 g/min.

4.18 Shifted true stress–strain curves for undrawn nylon 6 yarns obtained at different wind-up speeds, as indicated in the legend.

4000 m/min, on the other hand, the shifted true stress–strain curves coincide with the mastercurve only in the final strain-hardening regime. This result indicates a change in network topology with increasing wind-up speed. Based on a detailed structural investigation of nylon 6 yarns, it has been found that a transition in structure-formation mechanism occurs at wind-up speeds of about 3000 m/min, associated with the onset of spin line crystallization.
Below 3000 m/min, the as-spun nylon 6 is largely amorphous. It starts to develop crystallinity at above this speed. The crystals developed during spinning are primarily $\gamma$. At higher speed, more crystallization takes place. The partially crystallized filament yarn with $\gamma$ crystals, on conditioning, develops additional $\alpha$ crystallinity (post-crystallization). As the speed of spinning is increased, content of $\gamma$ crystals increases while that of $\alpha$ crystals decreases due to higher crystallization during spinning and limited post-crystallization. At spinning speeds of 5500 m/min or above, the sample has mostly $\gamma$ crystals. The effect of spinning speeds on birefringence profile and temperature profile of the spinning fiber is depicted in Figure 4.19.

The structure of the as-spun filaments depends markedly on spinning speed (take-up velocity) and on molecular weight. Although crystallinity and $\gamma$ fraction both increase with increasing take-up velocity, these quantities show an intricate dependence on molecular weight. The crystallinity at high take-up velocity (∼3000 m/min) decreases with increasing molecular weight. At low take-up velocity (∼1000 m/min) the opposite trend is observed, with the high-molecular-weight polymer having the higher crystallinity. This crossover in the behavior of the crystallinity is attributed to the fact that increasing either take-up velocity or molecular weight increases the crystallization rate but the final attainable crystallinity decreases with increasing molecular weight. The crystallization rate of the spun filaments increases with increase in molecular weight.

The tensile properties of the filaments are strong functions of both take-up velocity and molecular weight. The initial modulus and tensile strength increase and elongation to break decrease with increase of take-up velocity or molecular weight. The true strength at fracture is also found to increase with increase of molecular weight.

### 4.17 Drawing and heat setting

The as-spun fibers have poor elasticity (inability to recover) and they undergo plastic deformation on application of low levels of stress. This makes them unsuitable for use in applications including that of apparel. The polymer chains in the as-spun fibres are in partially folded conformation and can extend easily on application of stresses. Figure 4.20 shows a stress–strain plot of an as-spun fiber. The elastic region is small and is bounded by the yield point that occurs at a low stress and extension level. Following the yield point, the fiber shows deformation without much resistance till all the chains are unfolded and a new improved (more elastic) network of polymer is formed. In this region, the polymer chains assume new extended conformation and do not recover back to their original state when stresses are released. This region is known as region of ‘natural draw’ and the extensibility is called ‘natural draw ratio (NDR)’ of the fibers. This
region of easy extensibility must be removed to make the fibers show the behavior of increasing resistance with increasing extension.

Drawing is necessary to remove the region of natural draw of the fibers. When the fiber is drawn beyond the NDR, polymer chains form an elastic network with extended conformation in the direction of draw (i.e. along

4.19 On-line experimental measurements for nylon 6 melt spun with a mass throughput of 2.99 g/min. (a) Birefringence profiles; (b) temperature profiles.
the fiber axis). This gives fibers higher yield point, strength, initial modulus, and recovery compared to its as-spun form. Figure 4.20 also shows a typical stress–strain plot of a drawn fiber in comparison to that of an as-spun fiber. Maximum draw ratio (MDR), in contrast, is the maximum extensibility a fiber may be subjected to, before it breaks. The natural draw ratio in the as-spun fiber is a function of its spinning speed as discussed earlier.

The drawing is normally carried out by a cold drawing process where the fiber is drawn by forming a neck (also known as neck drawing). The location and stability of the neck is important in obtaining a uniformly drawn filament yarn. A snubbing pin may be used to help the formation of the neck or to stabilize it in a narrow zone. The drawing is carried out at just above the $T_g$ of the polymer. For apparel grade nylon 6, drawing is carried out at room temperature.

Drawing may be carried out in a single step called single stage drawing as in apparel grade yarn or in two or more steps called two stage drawing or multistage drawing, as in high performance tire yarns. There are certain advantages of drawing the material in more than one stage such as reduced defects in drawn fiber due to partial relaxation of critical stresses during the drawing process. Drawing is normally carried out as a separate process on a separate machine as described above; however, at times, drawing may be combined with spinning machine to provide in-line drawing.

As stated earlier, in drawing, polymer chains are unfolded and oriented by applying an external tensile stress in the direction of the fiber axis. The level of applied stress depends upon the state of as-spun fiber, the draw ratio, the draw rate, the draw temperature and the moisture content of the fiber. Filaments undergo flow deformation drawing when drawn at either...
a very low strain rate or at a high temperature. The filaments undergo a ductile failure when they are drawn at a high strain rate or at a low temperature. Therefore, in order to draw the filament in neck deformation, the rate and temperature of drawing must be optimized for a given fiber.

### 4.18 Mechanism of drawing in polyamides

The two theoretical schemes used for prediction of orientation from network deformation are (a) the affine and (b) pseudo-affine deformation mechanisms. In the affine scheme, network junctions are thought to be connected by flexible chains. Upon stretching, the network points are displaced in direct proportion to the macroscopic deformation. As a result, the rotatable ‘random links’ comprising the network chains will gradually adopt a more and more oriented configuration. In pseudo-affine deformation, on the other hand, the structural elements undergoing deformation are assumed to have no extensibility themselves but are rigid entities simply rotating in proportion to the macroscopic deformation of the sample.

The undrawn nylon 6 yarns are always partially crystalline, irrespective of the wind-up speed. The amorphous and crystalline orientation factors are plotted as a function of draw ratio in Figure 4.21. The crystalline orientation factor $f_c$ initially increases sharply, leveling off at higher draw ratio whereas $f_a$ initially rises very slowly becoming important only at the highest draw ratios. It appears that the crystalline phase responds to draw ratio in a pseudo-affine fashion whereas the amorphous phase follows a rubber-like, affine deformation. This result can be understood in

![Crystalline and amorphous orientation factors fc and fa of drawn yarns as a function of draw ratio determined from WAXS and solid-state NMR measurements.](image)
The crystalline structure of nylon 6 changes with the draw ratio at a given strain and temperature. This has been studied by measuring equatorial crystallinity index (ECI) with strain percentage as shown in Figure 4.22.41 This behaviour of ECI on strain percentage may be separated into three distinct regimes: regime I (0–15% strain); regime II (15–100% strain); and regime III (100% strain and greater). In regime I, the gamma ECI decreases while both the alpha and total ECI increase. The majority of this increase occurs after the first extension, from 0 to 5% strain. The increase of the alpha ECI is always greater in magnitude than the decrease of the gamma ECI in this region; so the net result, the total ECI, is increasing as well. In the second regime the a-phase, g-phase, and total ECI are all decreasing. Finally, in the third regime the \( \alpha \)-phase ECI remains constant and the \( \gamma \)-phase ECI drops to zero, thus forcing the total ECI to remain constant as well.

Since the increase of the a-phase ECI is greater than the decrease of the \( \gamma \)-phase ECI, a significant percentage of the \( \alpha \)-phase increase must be due to conversion of the amorphous phase as opposed to the \( \gamma \) to \( \alpha \) conversion. The decrease in the \( \gamma \)-phase ECI could be a result of two competing effects. Either the \( \gamma \)-phase is being converted to the \( \alpha \)-phase, as discussed above, or the \( \gamma \)-phase crystals are being destroyed during the drawing process.

### 4.19 Heat setting

Annealing tends to increase the amount of \( \alpha \)-phase mainly at the expense of the amorphous contribution. The nature of the \( \alpha \)-phases changes; it becomes monoclinic and increases in perfection as the severity of annealing increases.42 At low spin draw ratios there is very little \( \gamma \)-phase initially, but there is pseudohexagonal \( \alpha \)-phase present (\( \alpha^* \)). As noted by Illers5 and others43,44,45, there is no difficulty in transforming this paracrystalline pseudohexagonal \( \alpha \) to the monoclinic \( \alpha \) form. In the high spin draw ratio samples containing appreciable \( \gamma \)-phase, there is a decrease in the relative amount of \( \gamma \)-phase on annealing, but \( \gamma \)-phase is not eliminated by the most severe annealing treatment investigated (2 h in boiling aqueous 20\% formic acid solution). The low-SDR samples anneal more readily to the \( \alpha \)-phase.42
4.22 The equatorial crystallinity index (ECI) as a function of take-up speed and strain percentage of the: (a) $\alpha$-phase; (b) $\gamma$-phase; and (c) total ($\alpha + \gamma$); MPM = metres per minute.
4.20 References


5

Poly(lactic acid) fibers (PLA)

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5.1 Introduction

In a world that is becoming increasingly sensitive to the need to protect our environment, the ability to manufacture products from sustainable resources and which are fully compostable at the end of their useful life, is an exciting and attractive proposition. Poly(lactic acid) (PLA) is a linear aliphatic thermoplastic polyester derived from 100% renewable sources such as corn, and the polymer is compostable.\(^1\)\(^2\) However, most initial uses were limited to biomedical applications such as sutures\(^3\) and drug delivery systems\(^4\) due to availability and cost of manufacture. Over the past few years, NatureWorks LLC has developed large-scale operations for the economic production of PLA polymer used for packaging and fiber applications. It is important that PLA is used broadly in textile applications for several reasons. Polyesters currently used for apparel and related fiber applications, mainly poly(ethyleneterephthalate) (PET), account for over 40% of world textile consumption (second only to cotton) and their use is constantly increasing. Production of such polyesters consumes fossil fuel resources and disposal of the polymer adds to landfill sites as they are non-biodegradable and are not easily recycled. In contrast, PLA fiber is derived from annually renewable crops, it is 100% compostable and its life cycle potentially reduces the Earth’s carbon dioxide level. The recognition by the FTC in the USA and the EU commission that PLA fibers are a completely new generic class of synthetic fibers further reinforces the validity of this new approach to producing performance melt-spinnable fibers.

This chapter will review the chemistry of PLA and will discuss the commercial manufacturing process that confirms its position as a viable material for many applications in both the fibers and the plastics industries. It will review the various properties of the polymer and consider the fiber properties that make it attractive to the commercial sectors into which it is being developed today. Recognizing that this is very much a new polymer with its own characteristics and processing requirements, the current status
of the downstream technologies in these sectors will be appraised. The environmental benefits of PLA will be discussed and compared with the current petroleum-based polymers. The potential for further improvements in the environmental impact will be considered as the technologies for this new polymer are advanced.

5.2 Chemistry and manufacture of PLA polymer resin

Conventional synthetic polymers rely on reserves of oil and gas for their monomer source and energy to manufacture. These reserves of fossil fuel take millions of years to regenerate and are a declining resource. In contrast, the monomer used to manufacture poly(lactic acid) is obtained from annually renewable crops. Energy from the sun promotes photosynthesis within the plant cells; carbon dioxide and water from the atmosphere are converted into starch. This starch is readily extracted from plant matter and converted to a fermentable sugar (e.g. glucose) by enzymatic hydrolysis. The carbon and other elements in these natural sugars are then converted to lactic acid through fermentation (Fig. 5.1).5,6

Presently, the cheapest and most abundant source of sugar is dextrose (glucose) from corn. The land mass necessary for feedstock production is minimal. Producing 500 000 tonnes of PLA requires less than 0.5% of the annual US corn crop;7 since corn is a cheap dextrose source, the

![Diagram of the photosynthesis, starch formation, enzymatic hydrolysis, and fermentation processes to produce lactic acid from glucose.]

5.1 Production of lactic acid from renewable resources.
current feedstock supply is more than adequate to meet foreseeable demand. Furthermore, there are many other alternatives for the starch or sugar supply. As fermentation techniques improve, as PLA production improves, and if PLA production extends to other geographies, it is quite likely that other materials such as grass and even biomass could be used; there is no need to be reliant on food crops. PLA takes advantage of a biological system to do chemistry that traditional chemical techniques cannot.

5.2.1 Production of PLA

The polymer is formed either by (1) direct condensation of lactic acid or (2) via the cyclic intermediate dimer (lactide), through a ring-opening process (Fig. 5.2).²

**Polycondensation of lactic acid**

This process involves the removal of water by condensation and the use of solvent under high vacuum and temperature; the approach was originally used by Carothers who discovered PLA in 1932. With this route, only low to intermediate molecular weight polymers can be produced, mainly due to the difficulties of removing water and impurities. Other disadvantages of this route are the relatively large reactor required, the need for evaporation, recovery of the solvent and increased color and racemization. Most work has focused on the ring-opening polymerization, although Mitsui Toatsu Chemicals have patented an azeotropic distillation using a high boiling solvent to drive the removal of water in the direct esterification process to obtain high molecular weight PLA.
Ring-opening polymerization

This method is a better way to produce a high molecular weight polymer, and has now been adapted commercially due to advances in the fermentation of corn dextrose which have significantly reduced lactic acid production costs. The fermentation of sugar produces chiral lactic acid inexpensively in high yield. Chiral molecules exist as ‘mirror images’ or stereoisomers; lactic acid can exist as the L- or D-stereoisomer (Fig. 5.3). Chemically synthesized lactic acid gives the racemic mixture (50% D and 50% L), however, fermentation is very specific, allowing the production of essentially one major stereoisomer; fermentation derived lactic acid consists of 99.5% of the L-isomer and 0.5% of the D-isomer.

The process is based on removing water under milder conditions, without solvent, to produce a cyclic intermediate dimer, referred to as lactide. This monomer is readily purified under vacuum distillation. Ring-opening polymerization of the dimer is accomplished under heat, again without the need for solvent. By controlling the purity of the dimer, it is possible to produce a wide range of molecular weights. Production of the cyclic lactide dimer results in three potential forms: the D,D-lactide (called D-lactide), L,L-lactide (called L-lactide) and L,D- or D,L-lactide (called meso-lactide) (Fig. 5.4). Meso-lactide has different properties from D- and L-lactide; D- and L-lactide are optically active, meso- is not. Before polymerization, the lactide stream is split into a low D-lactide stream and a high D-/meso-lactide stream. Ring-opening polymerization of the optically active types of lactide can yield a ‘family’ of polymers with a range of molecular weights by varying the amount and the sequence of D-lactide in the polymer backbone. Polymers with high L-lactide levels can be used to produce crystalline

![L-lactic acid and D-lactic acid](image1)

5.3 The stereoisomers of lactic acid.

![L-lactide, Meso-lactide, D-lactide](image2)

5.4 Dimeric lactic isomers.

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polymers while the higher d-lactide materials (>15%) are more amorphous. By controlling the purity of the lactide it is possible to produce a wide range of molecular weights and by varying the amount and sequence of d-lactic units in the polymer backbone, the product properties can be changed. These changes impact melt behavior, thermal properties, barrier properties, and ductility.

Based on this lactide intermediate method, NatureWorks LLC has developed a patented, low cost continuous process for the production of lactic acid-based polymers. The process combines the substantial environmental and economic benefits of synthesizing both lactide and PLA in the melt rather than in solution and, for the first time, provides a commercially viable compostable commodity polymer made from annually renewable resources. The process starts with a continuous condensation reaction of aqueous lactic acid to produce low molecular weight PLA pre-polymer (Fig. 5.5).

Next, the pre-polymer is converted into a mixture of lactide stereoisomers using tin catalysis to enhance the rate and selectivity of the intramolecular cyclization reaction. The molten lactide mixture is then purified by vacuum distillation. Finally, PLA high polymer is produced using a tin-catalyzed, ring-opening lactide polymerization in the melt, completely eliminating the use of costly and environmentally unfriendly solvents. After the polymerization is complete, any remaining monomer is removed under vacuum and recycled to the beginning of the process (Fig. 5.6).
5.2.2 Catalytic polymerization of lactide

Many catalyst systems have been evaluated for the polymerization of lactide including complexes of aluminum, zinc, tin, and lanthanides. Metal alkoxides are the most common metal-containing species for the ring-opening polymerization of cyclic esters. Simple sodium, lithium, and potassium alkoxides are effective; however, the high basicity of these ionic species leads to side reactions such as epimerization of chiral centers in the polymer backbone. Alternatively, covalent metal alkoxides are much more selective and therefore widely used. Initiators such as Al-alkoxides, yttrium and lanthanide alkoxides, and recently iron alkoxides have been shown to give a controlled and living polymerization of lactides via a so-called coordination–insertion mechanism with ring opening of the lactide to add two lactic acid molecules to the growing end of the polymer chain. Depending on the catalyst system and reaction conditions, almost all conceivable mechanisms (cationic, anionic, coordination, etc.) have been proposed to explain the kinetics, side reactions, and nature of the end groups observed in lactide polymerization. Tin compounds, especially tin(II) bis-2-ethylhexanoic acid (Sn(Oct)₂), are preferred for the bulk polymerization of lactide due to their solubility in molten lactide, high catalytic activity, and low rate of racemization of the polymer; the mechanism is also via a coordination–insertion mechanism (Fig. 5.7). Conversions of >90% and less than 1% racemization can be obtained while providing polymer with high molecular weight. High molecular weight polymer, good reaction rate, and low levels of racemization are observed with Sn(Oct)₂ catalyzed polymerization of lactide. Typical conditions for polymerization are 180–210°C, Sn(Oct)₂ concentrations of 100–1000 ppm, and 2–5 hours to reach circa 95% conversion. The polymerization is first
order in both catalyst and lactide. Frequently, hydroxyl-containing initiators such as 1-octanol are used to both control molecular weight and accelerate the reaction.

Copolymers of lactide with other cyclic monomers such as \(\varepsilon\)-caprolactone\(^{17}\) can be prepared using similar reaction conditions (Fig. 5.8). These monomers can be used to prepare random copolymers or block polymers because of the end growth polymerization mechanism. Cyclic carbonates, epoxides and morphinediones have also been copolymerized with lactide.

### 5.3 PLA fiber properties

PLA fiber has a number of characteristics that are similar to many other thermoplastic fibers, such as controlled crimp, smooth surface and low moisture regain. One unique property in comparison is that it is the only melt-processable fiber from annually renewable natural resources. The physical properties and structure have been studied by several researchers,\(^{18}\) and these works confirmed that this polymer has significant commercial potential as a textile fiber. Its mechanical properties are considered to be broadly similar to those of conventional PET,\(^{19}\) and, probably due to its lower melting and softening temperatures, comparisons to polypropylene are also appropriate.\(^{20}\) A résumé of the properties is given, although further detail about specific properties will be covered, as appropriate in Section 5.4, ‘PLA Applications’:
• **Appearance.** Fibers are generally circular in cross-section and have a smooth surface.

• **Density.** The specific gravity is 1.25 g cm\(^{-3}\), lower than natural fibers and PET.

• **Refractive index.** The refractive index of 1.35–1.45 is lower than PET (1.54). Trilobal and other shapes can be made, and give improved anti-soiling characteristics.

• **Thermal properties.** PLA is a stiff polymer at room temperature. The glass transition temperature (T\(_g\)) is typically between 55–65°C. The melting temperature (T\(_m\)) of PLA containing either the L- or D-isomeric form alone, is between 160–170°C. The DSC scans for PLA and PET are shown\(^\ref{fig:dsc-pla-pet}\) (Fig. 5.9). It can be seen that PLA exhibits an endothermic peak (T\(_m\)) at approximately 166°C, whereas the T\(_m\) of PET is approximately 254°C. This low melting point compared to PET underlines one of the main restrictions for PLA in developing suitable applications. However, as mentioned before, the properties of PLA can be modified by adjusting the ratio and the distribution of the D- and L-isomers in the polymer chain, and melting points as low as 130°C and as high as 220°C have been obtained.

• **Crimp.** PLA can achieve good degree of crimp and good retention level through processing.

• **Fiber types.** Both filament yarns and spun yarns can be made, as with PET.

• **Tenacity.** The tenacity at break (32–36 cN tex\(^{-1}\)) is higher than for natural fibers although, of course, it can be varied according to the

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5.9 DSC scans of PET and PLA.
degree of drawing that is applied to the undrawn yarn. It is relatively unaffected by changes in humidity at ambient temperature, though as with other manufactured fibers there is a small but measurable increase in elongation. As the temperature is increased the tenacity does reduce quite quickly with a concomitant increase in fiber extension, a feature commonly found in synthetic fibers.

- Tensile properties. The tensile properties of PLA fiber as used in staple form for textile processing are shown in Fig. 5.10. Clearly they are very different from those of high tenacity polyester and more akin to wool with a high fiber extension when stressed and relatively low final tenacity. The initial modulus (at 2% extension) is very similar to many other textile fibers, but the yield point is very marked, the fibers (and spun yarns) stretching very easily once past this point. A consequence, however, of the high elongation is that the work of rupture is relatively high giving yarns and fabrics an acceptable performance in commercial use. Elastic recovery is affected by the yield point and is particularly good at low strains. At 2% strain, the recovery is 99.2% \(+/-0.75\%\), and 92.6% \(+/-1.60\%\) at 5% strain, higher than for most other fibers. The unusual tensile properties could be expected to have an influence on some of the commercial applications in fiber blend developments. For example, intimate spun yarn blends with cotton (and lyocell) are relatively weak due to the contrast of fiber properties, and this might possibly limit applications to knitted fabrics. However, wool blends well with PLA, and with their load-elongation curves being very similar, this would enable the full properties of both fibers to be exploited.
• **Moisture regain.** At 0.4–0.6%, PLA has extremely low moisture regain, much lower than natural fibers and slightly higher than polyester.

• **Flammability.** Although PLA is not a non-flammable polymer, the fiber has good self-extinguishing characteristics; it burns for two minutes after a flame is removed, and burns with a white and a low smoke generation. PLA also has a higher LOI (limiting oxygen index) compared to most other fibers, meaning that it is more difficult to ignite as it requires a greater oxygen level. Table 5.1 compares PLA with standard PET.

• **UV resistance.** Unlike other synthetic fibers, PLA does not absorb light in the visible region of the spectrum; this leads to very low strength loss compared to petroleum-based fibers when exposed to ultraviolet light.

• **Moisture transport.** PLA shows excellent wicking ability. This property and the additional properties of fast water spreading and rapid drying capability give the fiber a very positive inherent moisture management characteristic.

• **Biological resistance.** Although PLA fibers are not inherently ‘antimicrobial’ without suitable after-finish treatment, they do not provide a microbial food source. In addition, testing by Odor Science and Engineering showed that PLA fiber-based fabrics outperformed PET-based fabrics for low odor retention.22

• **Chemical resistance.** As PLA is a linear aliphatic fiber, its resistance to hydrolysis is therefore relatively poor. This feature means that care must be taken in dyeing and finishing of the fiber.

• **Solubility.** With regard to other chemicals PLA has limited solubility and is unaffected by dry-cleaning solvents for example.

### 5.4 Applications

The ease of melt processing, coupled with the unique property spectrum and renewable resource origin, has led to PLA fibers finding increasing
acceptance across a variety of commercial sectors. It would also seem that with the estimated global increase in fiber demand, there is an opportunity for viable alternative materials. The total fiber business in 2002 was estimated to be some 55 million tonnes, and projections suggest that this could increase to 83 million tonnes by 2015. Particularly relevant within these figures is the growth of man-made fibers from 33 million tonnes in 2002 to 57 million tonnes by 2015, and that the major fibers creating this increase will be synthetic materials, mainly PET.23

With approximately 50% of the fibers in the market today being synthetic materials from petrochemicals, it is clear that without synthetic fibers there would be insufficient fiber materials to meet the demands, particularly as natural fibers alone (mainly cotton) could not cover this. It is considered that cotton is already at near peak production, so as the world population and the fiber requirements increase, the opportunity to use such a product as PLA is both logical and in line with our need to address the environmental problems we face.

5.4.1 Apparel

The apparel fiber business in 2002 was estimated at approximately 30 million tonnes (Table 5.2) and projections for 2015 indicate that it will grow to 42 million tonnes. The sector is dominated by two fibers, cotton and PET, and as the total requirement increases, it is PET that is projected to become the majority material. Apparel is by some way the largest sector in the global fiber business, and is a highly technical, fast changing, and demanding industry in terms of design, fashion, color, aesthetic, and performance. However, despite the steady growth in the apparel fiber market size, there has been a reduction in the price points, therefore creating a real pressure on costs through the whole supply chain. In many respects,

<table>
<thead>
<tr>
<th>Fiber type</th>
<th>Million tones</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cotton</td>
<td>13.0</td>
<td>43</td>
</tr>
<tr>
<td>Wool</td>
<td>1.3</td>
<td>4</td>
</tr>
<tr>
<td>Polyester</td>
<td>10.2</td>
<td>34</td>
</tr>
<tr>
<td>Acrylic</td>
<td>2.0</td>
<td>7</td>
</tr>
<tr>
<td>Cellulosic</td>
<td>1.8</td>
<td>6</td>
</tr>
<tr>
<td>Nylon</td>
<td>1.2</td>
<td>4</td>
</tr>
<tr>
<td>Others</td>
<td>0.5</td>
<td>2</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>30.0</strong></td>
<td><strong>100</strong></td>
</tr>
</tbody>
</table>

Table 5.2 Apparel mill consumption by fiber type 2002
it would seem that the existing range of fibers could meet all the possible consumer needs, either by virtue of their inherent properties, or with some chemical or physical processing technique within the downstream processing. For a new fiber, the entry into the apparel market could therefore be seen as both a very exciting challenge, but also a very difficult one. Against this backdrop, the real opportunity for PLA lies in the combination of both the fiber’s inherent performance properties and the positive environmental advantages of being sourced from annually renewable natural resources, which, at the end of the product life, can be easily composted or recycled.

Kanebo, Inc. introduced a PLA fiber under the trade name Lactron™ fiber at the February 1998 Nagano Winter Olympics, under the theme of ‘Fashion for the Earth’. Kanebo exhibited several garments from PLA or PLA/natural fiber blends. More recently, in 2003, Cargill Dow LLC (now NatureWorks LLC) announced their PLA fiber brand Ingeo™ (ingredient from the earth), and this is now being adopted across a wide range of garment categories globally by leading garment brands.

It is one of the features of PLA that it can be produced as both filament and spun yarns. Fabrics produced from spun yarn have a ‘natural’ hand and are considered to feel similar to cotton in this respect. Fabrics from filament yarns have a cool and soft hand and exhibit a high fluidity or drape with a degree of elasticity. NatureWorks LLC product development suggests, for example, that a 1.2 dpf PLA achieves the softness of a micro-denier PET (e.g. 0.7 dpf).

With the opportunity to create a range of fabrics with attractive aesthetic properties, there are a number of performance properties of PLA that are significant to its positioning and progress in the apparel sector:

- The moisture management properties of good wicking, faster moisture spreading and drying mean that garments are comfortable. This is an important consideration for next-to-skin garments and particularly in the sportswear market. In terms of comfort under normal and active wear conditions, independent laboratory testing by the Hohenstein Research Institute demonstrated that PLA fibers perform better than PET and cotton, either when combined with cotton or as 100% PLA fabric. This testing included a series of different tests, including thermal insulation, breathability, water vapor transport, and buffering capacity to liquid sweat (several tests in this series). The results of the Hohenstein testing concluded that wearers of PLA/cotton fabric would experience improved physiological comfort versus equivalent PET/cotton fabric.24
- The elastic recovery and crimp retention properties provide excellent shape retention and crease resistance.
Table 5.3 AATCC Test Method 61-1994 (35% PLA/65% cotton blend knitted shirt, simulates five washings)

<table>
<thead>
<tr>
<th>Simulated conditions</th>
<th>AATCC Test</th>
<th>Burst strength (psi)</th>
<th>% Dimensional change (width/length)</th>
<th>M_n</th>
<th>M_w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td></td>
<td>83</td>
<td>–</td>
<td>57694</td>
<td>117970</td>
</tr>
<tr>
<td>Cold hand wash (40°C)</td>
<td></td>
<td>79</td>
<td>0/−3.82</td>
<td>56343</td>
<td>107835</td>
</tr>
<tr>
<td>Same, no bleach</td>
<td>1A</td>
<td>82</td>
<td>0/−3.13</td>
<td>52123</td>
<td>108115</td>
</tr>
<tr>
<td>Cold machine wash</td>
<td>–</td>
<td>75</td>
<td>0/+4.17</td>
<td>56281</td>
<td>111206</td>
</tr>
<tr>
<td>Same, no bleach</td>
<td>–</td>
<td>78</td>
<td>0/−3.82</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Warm machine wash</td>
<td>5A</td>
<td>74</td>
<td>+6.25/−7.98</td>
<td>57190</td>
<td>112036</td>
</tr>
<tr>
<td>Same, no bleach</td>
<td>2A</td>
<td>78</td>
<td>+7.64/−7.98</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hot machine wash</td>
<td>4A</td>
<td>74</td>
<td>+2.00/−6.25</td>
<td>58005</td>
<td>112510</td>
</tr>
<tr>
<td>Same, no bleach</td>
<td>3A</td>
<td>76</td>
<td>+2.00/−6.25</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Thermosetting capability of the fiber provides for controlled fabric stability, with garments having a low shrinkage through repeated washings.
- The flammability properties described earlier show the potential in specific segments of the apparel market. PLA fabrics with no flame retardant treatments have passed the US tests 16 CFR 1610, and have also achieved the standards specific for children’s sleepwear, 16 CFR 1615 and 16 CFR 1616.
- High resiliency and lower specific gravity than natural fibers can give fabrics a light and lofty feel. These properties have been used in garments to provide an effective wadded layer, e.g. ski jackets.
- The after-care properties of garments in washing are very positive. There is no damage in repeated laundering of PLA fabrics: testing has been carried out under simulated conditions in accordance with AATCC standards, with no degradation observed (Table 5.3). Also, the fabric appearance remains extremely good after washing, without creasing and very clean surface. It would seem that any fiber work-up, which is quite normal under such conditions, is removed during the washing cycles.

Additional to the above properties that could all be described as positive attributes, there are certain factors in this relatively early stage of technical and commercial development that are somewhat restrictive to the development across a full apparel spectrum:
• The melting point of the yarns that are commercially available today is relatively low at 170°C. This does cause limitations in some of the downstream processing technologies, but the main concern is for the consumer after-care of garments. Garment pressing and ironing temperatures have to be lower than the popular fibers of cotton and PET, and despite the appropriate care labeling instructions being used, it is a fact that consumers often disregard these.

• Hydrolysis degradation of the polymer can occur, particularly under combined aqueous high temperature and alkaline conditions; the degree of hydrolysis is influenced by the time, temperature and pH. This is of particular significance in the dyeing and finishing processes, as it will cause a reduction in the molecular weight of the polymer and therefore the strength of a yarn or fabric, if the appropriate finishing conditions are not observed.

Through all of the manufacturing stages of the supply chain – spinning, fabric formation, dyeing and finishing, and garment making – the existing machinery can be used. There is no need for any specialized capital investment when processing PLA. In many of the downstream processing technologies, PLA processes and products are compared and are comparable to PET; both are melt spun, both are available in filament and staple form, weaving and knitting set-up conditions are similar, fabrics can be heat treated to give dimensional stability, and they are both dyed with disperse dyes. However, both fibers are unique and they do behave differently. It is therefore important that processors acknowledge this fact, and that the operating details are refined accordingly.

Yarn spinning

There is now a wide selection of commercial yarns being produced. In spun yarns, commercial products currently span the range Ne 5 to Ne 60 while in filament production, typical products such as dtex 70–68, dtex 150–72, are being produced. For both spun and filament yarns spinners have confirmed the processability of the material. An interesting feature of PLA is that processing temperatures are more typical of polyolefins (approximately 240°C), although the properties more typically combine many of the features found in rayon, nylons, acrylics and polyesters. PLA pellets are supplied with a maximum moisture content of 400 ppm. It is important to reduce the moisture content to 100 ppm during processing to prevent hydrolytic degradation, similar to PET and nylon products. For short staple yarns, because the characteristics of PLA are comparable to other thermoplastic fibers, the processing conditions are similar to those for PET. For long staple yarns also, processing can be carried out on conventional
long staple equipment as used for thermoplastic fibers in blend with wool.

**Fabric formation**

There are no special requirements for knitting, and machine settings are comparable with those for PET. In weaving, the additional points for attention would be: if a size is being applied, to use a PVA or water-soluble size to avoid any need for strong alkali desizing, and to minimize tensions due to the high fiber extension. Woven filament fabrics give a very soft hand, and have a high fluidity/drape compared to PET.

**Garment making**

The low melting point may mean that precautions are taken at those stages where fiber–metal friction could generate sufficient heat, namely the fabric cutting and the garment panel sewing operations. Care is also needed at the final pressing stage.

**Dyeing and finishing**

As is usual with any new fiber, the coloration and wet processing technologies demand significant attention. Color is so important, not only because it is the prime driver when purchasing a new garment, but also because the various applications and subsequent finishing treatments have the most chemical and physical effect on any fiber or polymer. Similar to PET, PLA is dyed with disperse dyes. However, dye selection is most important, as the individual dye behavior is quite different from dyeing on PET. In general terms, dyes show their maximum absorption at a shorter wavelength than on PET and tend to look brighter\(^2^5\) (Fig. 5.11). Dyes also show a much greater variation in exhaustion levels; Yang and Huda\(^2^6\) studied the exhaustion of ten disperse dyes on PLA and PET fabrics and found that the percentage dye exhaustion of all the dyes was lower on PLA than on PET. However, the color yield of the dyed PLA was higher than on PET because of the lower reflectance of PLA.

Several of the major dye manufacturers have studied their ranges of disperse dyes to provide the most appropriate selection for applying to PLA. DyStar GmbH & Co. have worked closely with NatureWorks LLC to further the commercial dyeing technology for PLA and have recently provided details for dyeing both 100% PLA as well as blends with cotton and wool.\(^2^7\) The optimum dyeing conditions recommended by DyStar for dyeing PLA is 110°C for 30 minutes at pH 5. Practical experience has shown
that the use of higher temperatures or longer times of dyeing can cause degradation of the polymer.

One of the observations in the dyeing of PLA is that obtaining dark shades is more problematic, compared to PET. A reason for this is attributable to the lower exhaustion levels, although, of course, dye selection has to balance many other factors including fastness requirements, reproducibility, and levelness.

There are indications that the color exhaustion and color yields can be increased by modification of the basic PLA polymer, altering the proportions of the D- and L-isomers, and thereby changing the amorphous/crystalline ratios. Higher D-levels have more amorphous and less crystalline regions and allow for increased dye exhaustions. Comparison of PLA fabrics with varying D-isomer content revealed differences in the enthalpy of fusion, and, hence, percentage crystallinity; high D-fibers have more amorphous and less crystalline regions in the polymer, with respect to low D-fibers. High D-fabrics display greater dye exhaustion and color strength with respect to low D-fabrics in all dyes and all concentrations as a result of the greater number of amorphous regions in high D-fibers with respect to low D-fibers. In application of a dye mixture for a black shade high D-fibers are able to be dyed to an excellent black shade, whereas low D-fibers appear very brown, due to less exhaustion of the blue component of the mix. In terms of wash fastness, there is very little difference between high D-fibers and low D-fibers; this is because the glass transition temperature for both fibers is very similar.

Color fastness figures tend to be slightly lower than on PET. One reason for lower wet fastness is believed to be due to more movement of dye to the surface by thermomigration, during post-heat treatments. It may also

![K/S curves of C.I Disperse Red 60 on dyed PLA and PET.](image)

**5.11 K/S curves of C.I Disperse Red 60 on dyed PLA and PET.**
be that the lower thermosetting temperature for PLA (130°C) compared to PET (190°C) means less sublimation of the dye from the surface.²⁹

With the introduction of PLA into apparel, developments and commercial adoptions have included fabrics made from 100% PLA as well as in blends with other fibers. The main blends are either with cellulosic fibers (cotton, lyocell) or wool. Apart from any aesthetic or performance benefits, such blends also have the feature of being biodegradable compared to their PET counterparts.

The wet processing of cellulosic blends need to be adapted to recognize the sensitivity of PLA to alkali treatments. This causes some limitations, as bleaching and dyeing systems for cellulosic fibers generally use alkaline processes. However, the potential significance of this blend has been recognized, and methods are available for all stages of wet processing. These include the use of neutral bleaching systems based on TAED (tetraacetyl-ethylenediamine), and direct dyes, as well as the more conventional alkaline bleaches and selected reactive dyeing systems. A detailed study into the effect of the various wet processes on the molecular weight and physical strength of PLA was carried out at UMIST, and this confirmed the feasibility for suitable processing.³⁰,³¹ Dyeing of PLA/wool blends does not present such a problem, as both fibers have the same characteristic with regard to alkali. Indeed, with their similar stress–strain profile, there would seem to be some benefits compared to PET/wool.

There is much still to learn about the dyeing of PLA but it would seem that processes are available to achieve commercial viability. In real terms, it is not that long ago that fibers such as lyocell and indeed PET were introduced to the textile world, with many technical difficulties in their processing. These are now well accepted fibers and it will be interesting to watch as PLA follows a similar course.

### 5.4.2 Homeware

Typical products encompassed in this segment range from pillows, duvets, blankets, mattress pads, carpet tiles, office panel fabrics, drapes to bonded fiber products such as mattresses. As in the other sectors, the unique origin of these fibers, being derived from natural sugars obtained from annually renewable resources, coupled with the performance benefits, is proving to have a strong appeal to consumers in various geographies. The resistance to UV and low flammability, low smoke generation and low toxic gas on burning are attractive properties for this market segment, which differentiate PLA fabrics manufactured from conventional petrochemical-based synthetics.

Superior resilience found in crimped staple fiber products such as fiber-fill, coupled with the natural wicking performance of the fiber are added
features, which further enhance the scope of opportunities for PLA. Since this moisture wicking behavior is inherent to the fiber and achieved without the use of finishes this behavior should not decrease over time. Independent testing indicates the wicking behavior of untreated PLA fibers is superior to either untreated or treated polyester fibers. Laboratory UV resistance testing using a Xenon Arc indicates that in comparison with polyester and acrylic fibers, PLA fabrics have superior strength retention than polyester and far superior resistance to discoloration than acrylcs.

Independent testing by the Hohenstein Institute shows that PLA fibers when used in duvets provide a better microclimate between the body and the duvet. The superior wicking properties of PLA fibers, compared with polyester, results in the dynamic adjustment of the moisture level by dissipating moisture as the humidity level changes. In pillows, testing has shown that PLA fibers offer outstanding filling power and resilience after three years of simulated usage.

Flammability testing indicates a Limiting Oxygen Index value for PLA fibers of 24–28 when tested according to ASTM D2863. This is superior to untreated PET fibers, cotton, rayon and acrylics and equivalent to natural protein-based fibers such as wool. In addition, flame propagation and time to extinguish are properties of considerable interest in the furnishings segment. Fabrics made from untreated PLA fibers show far shorter burn times than cotton and polyester fibers. Untreated PLA fibers meet the test criteria for UNI 8456(1986), UNI 9174(1987) and UNI 9174/A1 (1996) that measure the time to self-extinguish (after the fabric is removed from the heat source). This test also measures the rate of flame propagation, char length, and time of flaming of any dripping materials to extinguish. The results enable 100% PLA fabrics to be classified as Category 1 as required by the Italian Drapery Industry.

Fabrics made from PLA fibers have also passed flammability tests such as NFPA 701, which covers the fire safety requirements for textiles used in interior furnishings for public occupancy buildings. Typical products include drapery, cubicle curtains, wall covering and protective outdoor coverings such as tarpaulins and tents.

Finally, a prototype mattress made from 100% PLA fibers successfully passed Cal 129 testing for furniture flammability. All these tests were conducted on 100% PLA without any flame retardant additives. However, individual results will depend upon fabric construction and the use of finishes or dyestuffs.

The unique resistance to staining and soiling of PLA Bulk Continuous Filament (BCF) has led to significant interest in PLA fibers for use in carpets. Presently, PLA fibers are used in carpet mats for the hybrid Toyota Prius in Japan as well as in carpet tiles for domestic and institutional markets.
As described earlier, PLA bicomponent technology utilizes the unique polymer properties induced by the two optically active forms of the lactic acid monomer. By controlling the ratio and distribution of the \(d\)- and \(l\)-isomers in the polymer chain, it is possible to induce different crystalline melting points during melt processing; this feature offers distinct benefits particularly in the binder fiber area. Bicomponent fibers with a sheath melting point of 60°C (amorphous) up to a crystalline melting point of 175°C can be produced (Fig. 5.12). As well as bonding to other PLA fibers or petroleum-based synthetics, PLA bicomponent binder fibers are increasingly being used to bond natural fibers such as jute and hemp in automotive applications for spare wheel covers and door panels. Bicomponent PLA technology is also finding applications in self-crimping and micro denier technologies. Technology growth areas are in sheath/core, side by side, segmented pie and islands in the sea structures.

Overall, PLA binder fibers can replace existing synthetic polymers where renewable resource is a benefit or where additional performance such as controlled temperature bonding, controlled shrinkage or lower temperature processing is required. In addition, improved or reduced adhesion and alternative approaches to soluble/non-soluble island/sea combinations are all possible and under development.
5.4.3 Nonwovens

Nonwoven products are a major application segment offering great potential for the unique benefits of PLA fibers. Outside of the fiberfill products the major markets are in spun bond, industrial and household wipes, hygiene and filtration areas. Spun bond PLA products can be produced on typical polyester spun bond lines in a variety of fabric weights. Filament velocities of 3500 m min\(^{-1}\) and above are needed to produce fabrics with the required low shrinkage performance. Key applications are in carpet backing, hygiene, and compostable geotextiles for soil erosion control and plant as well as crop protection. Under the correct conditions of temperature and humidity PLA fabrics are completely compostable and return naturally to the soil releasing carbon dioxide and water. Under normal storage conditions and use, however, the fabrics are durable enough to meet the various market and supply chain requirements.

*Industrial/household wipes*

In recent years there has been a significant growth in the wipes market, and in particular the wet wipes segment across a variety of applications. Most wet wipes comprise a blend of cellulose, viscose or rayon with a synthetic fiber such as polyester or polypropylene. Up to 50% of the wipe comprises these synthetic fibers. Recent surveys indicate a 6.5% annual growth in personal care wipes versus 2.5% in baby wipes in the US.\(^{37}\) Similar trends are being seen in Europe and Japan. New product introductions in personal care and household cleaning markets are occurring rapidly. Typical new trends and applications for wet wipes include:

- Feminine hygiene, facial cleansing, hemorrhoid treatment, etc.
- Functional treatments (antimicrobial, cleaners, abrasives, etc.).
- Decorative patterns.
- Continued push toward cloth-like aesthetics.
- Environmentally friendly materials and processes.

Spun laced investments have recently been announced by Jacob Holm, Spuntech, Orlandi, and Green Bay Nonwovens. In addition, Japan has large spun lace capacity with markets that are seeking environmental solutions. Wipes converters globally are expanding capabilities and expediting line extensions. PLA fibers show superior wicking performance when compared with the petroleum-based synthetics used in these applications. This inherent property leads to increased rates of liquid absorption,\(^{32}\) in addition the appeal of an all-natural-based wipe has demonstrated strong consumer pull. The wipes market is, however, extremely price sensitive; recent price fluctuations in the price of oil coupled with the advances made in

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PLA fiber and resin production means that pricing differences between the petrochemical-based and PLA fibers continues to narrow.

**Hygiene**

PLA fibers, because of their natural wicking properties, are finding utility in the diaper and feminine hygiene markets, both in spun bond top sheets and acquisition/distribution layers. Again the ability to replace petroleum-based fibers with enhanced performance with a natural-based fiber has strong consumer appeal. An additional benefit is that since PLA fibers are fully compostable the products can be disposed of by composting if the infrastructure for disposal is in place.

**Filtration and separation**

Two areas of potential applicability of PLA polymers have appeared in the industry in recent years.

- Triboelectric media to improve filtration efficiency.
- Filters, for example in the automotive and chemical industries, as well as single use applications.

The unique combination of properties displayed by PLA fibers offers promise in these areas. Untreated PLA fibers are repelled by glass surfaces and are attracted to polypropylene, indicating that they are somewhat electropositive. Disposable filters are also an area of interest for PLA fibers.

**5.4.4 Medical applications**

Textile fibers can be used to cultivate different human organs. The process involves culturing and growing living cells, taken from human organs, on a textile scaffold, to the desired 2-dimensional and/or 3-dimensional shapes. The scaffold is made from biodegradable and resorbable fibers which are in turn produced from biocompatible and degradable polymers. The major bioresorbable fibers used in implants are PLA and PGA (polyglycolic acid). They can either be used as a single polymer or by blending a copolymer of PLA and PGA. Varying the proportions of PLA and PGA alters the degradation rate and strength retention time of the fiber. These properties can therefore be varied in this way according to the requirements of specific medical applications. During the process of degradation, fibrous connective tissues replace the degrading implant. The key advantage is that no further surgery is required to remove the products since they slowly degrade in the body without any side affects. The US Food and Drug Administration (FDA) has approved the use of PLA for certain human
clinical applications. Also, PLA-based materials have been used for bone support splints.

5.4.5 PLA as a plastic

Although it is outside the scope of this chapter, it is important to recognize the tremendous scope for PLA as a plastic in several applications. The material has significant potential use in fresh food rigid packaging, bottles for beverages like milk, juice and non-carbonated water. PLC can also be made into an ideal film for use as labels, wrappers and windows.

5.5 Environmental sustainability

The previous sections of this chapter have examined the manufacturing of PLA, its various properties as a polymer and a fiber, and considered the potential for its use in a range of commercial applications. Additionally, the environmental aspects for producing and using PLA products must be appraised in any discussion about its significance as a sustainable commercial polymer material.

Environmental sustainability is about making products that serve useful market and social functions with lower detrimental environmental impact than the currently available alternatives. The case for PLA in this respect would seem to be very strong and needs to be appraised in any discussion about its potential as a significant polymer material. The ideal environmentally sustainable product provides equivalent function as the product it replaces and is available at competitive costs. It is made from renewable resources, can itself be constantly renewed without degradation in quality or performance, and has a minimum environmental impact. Such a product is made using only substances known to be safe for both humans and the environment. Ideally the life cycle of the sustainable product is in balance with the surrounding ecosystem.

5.5.1 Polymer processing and environmental measures

According to Vink et al., there are probably three items that are generally considered to have an increasing global importance with regard to environmental concerns: fossil energy use; greenhouse gas emissions; and water use.

Fossil energy use

The existing range of petrochemical-based plastics is diverse, specialized and mature, so that precise and exact comparisons with PLA, a single
5.13 Fossil energy requirement for some petroleum-based polymers and polylactide. The cross-hashed part of the bars represents the fossil energy used as chemical feedstock (the fossil resource to build the polymer chain). The solid part of each bar represents the gross fossil energy use for the fuels and operations supplies used to drive the production processes. PC = Polycarbonate; HIPS = High impact polystyrene; GPPS = General purpose polystyrene; LDPE = Low density polyethylene; PET SSP = Polyethylene terephthalate solid state polymerization; (bottle grade); PET AM = Polyethylene terephthalate Amorphous (fibers and film grade); PP = Polypropylene; PLA1 = Polylactide (first generation); PLA B/WP (Polylactide, biomass/windpower scenario).

product performing multiple functions, are difficult, especially considering the great number of impact categories compared. Figure 5.13 plots the fossil energy requirement for these products. Data for the petroleum-based polymers was supplied by the Association of Plastics Manufacturers in Europe (APME). The data is valid for the polymers as produced in Europe. A key finding of the analysis is that the first generation polylactide production system (PLA1) uses 25 to 55% less fossil energy than the petroleum-based polymers.\textsuperscript{38,39} Process improvements are targeted by NatureWorks LLC for the near future involving the use of biomass (B) and wind power (WP) as energy sources in the PLA production process (PLA B/WP), and with these improvements the use of fossil energy can be reduced by more than 90% compared to any of the petroleum-based polymers being replaced. This also will give a significant reduction in fossil energy related air and water emissions. This comparison represents the outstanding
potential for environmental benefits for polymers made from renewable resources.

It needs to be recognized that the data for PLA1 and PLA B/WP represent engineering estimates. In addition, there is good reason to expect improvements in the actual performance versus the estimates. Despite years of development work, the commercial manufacturing process for PLA is in its infancy. If the experience from petrochemical-based polymers offers any instruction, it is that process improvements implemented in the early years of a technology typically lead to substantial cost improvements. This is because the pursuit of cost improvements for competitive reasons often targets energy use due to its relatively high contribution to overall material costs. For example, through work on biocatalyst and lactic acid manufacturing process, NatureWorks LLC expects to achieve improvements that should further improve the performance of their production plant and simultaneously reduce energy demand. There is therefore good reason to expect a performance improvement trajectory for PLA1 that mirrors the experience from the current incumbent materials.

**Global climate change**

Global climate change has been identified as perhaps the most important environmental issue of this century. Greenhouse gas emissions are not exactly the same as combusted fossil fuel emissions, because several non-combustion gases can contribute to global climate change as well. For example, methane (CH₄) is a potent greenhouse gas that can emanate from natural gas system leaks, decomposition of biological materials, and chemical/industrial processes. However, greenhouse gas emissions are closely correlated to fossil fuel emissions because combustion of fossil fuels is the source of most anthropogenic greenhouse gases. NatureWorks LLC has undertaken a comparison of the contributions to global climate change from a range of petrochemical-based polymers as well as the two PLA cases described above. This comparison is depicted in Fig. 5.14. In conducting this analysis, Vink et al. relied upon the 100-year time horizon Global Warming Potentials for greenhouse gases, a time period generally accepted as the mean atmospheric residence time for the most volumetrically significant greenhouse gas, carbon dioxide. A check of the data revealed that use of the 20- and 500-year time horizons generates the same ranking among the products studied. According to the Intergovernmental Panel on Climate Change (IPCC) the relative global warming potentials of the three largest (volumetric) greenhouse gases are: CO₂-1, CH₄-21, and N₂O-310. These factors were used in NatureWorks LLC’s analysis. As in the comparison of fossil energy use, the analysis compares conventional polymers with PLA from cradle to pellet (from raw materials to the point
where the product is ready for shipment to a converter or fabricator). All emissions values were converted to CO₂ equivalents in order to facilitate comparison.

The analysis demonstrates that the PLA1 production process enjoys a substantial advantage over most polymers, and is comparable to several others. Even more exciting are the greenhouse benefits that derive from the transition to biomass feedstocks and reliance on wind energy for the balance of plant energy requirements. The utilization of the lignin fraction of lignocellulosic feedstocks for process heat generation ‘closes the loop’ on carbon related to energy generation, and in combination with other factors yields a negative greenhouse gas impact for PLA pellets. A most appealing result of the use of agricultural feedstocks for the PLA polymer production and most of the process energy requirement means that customers using PLA cannot only use PLA as a product, but as a component of their greenhouse gas reduction strategies.

Life cycle assessment reveals that no petroleum-derived polymer can rival the greenhouse gas sink effect of the improved PLA process. Although disposal of PLA products – whether by combustion, composting or other conventional means – results in a return of carbon dioxide to the atmosphere, this advantage survives.

**Water use**

Vink et al. also studied the water use. Figure 5.15 gives the gross water use of the traditional polymers and the two PLA cases (PLA1 and PLA B/WP) as described above. The gross water use is the sum of public supply, river, canal, sea and well water and used as cooling water, process water and irrigation water. Despite the use of irrigation water during corn growing...
and the two water-based processes (dextrose and lactic acid production) the total amount of water required is competitive with the best performing petrochemical polymers.

5.5.2 Disposal options

The most common waste management options for the fossil fuel-based polymers are incineration, landfill and mechanical recycling. In addition to these traditional processing routes, the PLA waste streams can also be processed using composting, chemical recycling and anaerobic digestion.

Composting

Composting is a beneficial waste management system, particularly where landfill sites are limited, and in more densely populated locations. It does require an appropriate infrastructure to be set up, but progress is being made, particularly in parts of Western Europe. Composting is a method of waste disposal that allows organic materials to be recycled into a product that can be used as a valuable soil amendment. The primary mechanism of degradation of PLA is hydrolysis, catalyzed by temperature, followed by bacterial attack on the fragmented residues. In composting, the moisture and the heat in the compost pile attacks the PLA polymer chains and splits them apart, creating smaller polymer fragments, and finally, lactic acid. Microorganisms, found in active compost piles, consume the smaller polymer fragments and lactic acid as energy source. Since lactic acid is
widely found in nature, a large number of naturally occurring organisms metabolize lactic acid. At a minimum, bacteria and fungi are involved in PLA degradation. The end result of the process is carbon dioxide, water and some humus. In summary, via composting, the carbon dioxide which has been harnessed during corn growing flows back into the atmosphere and the short cycle carbon dioxide loop has been closed. The degradation process is temperature and humidity dependent. PLA is compostable at domestic composting piles since the minimum required conditions are typically not present. A typical degradation curve of PLA under composting conditions is shown in Fig. 5.16.

Chemical recycling

Vink et al. describe the possibilities of chemical recycling as a promising future alternative waste disposal route. The PLA polyester polymer is formed from reversible polycondensation reactions and can be depolymerized by hydrolysis. This equilibrium results in recycling advantages for polyesters such as PLA. Manufacturing waste, converter waste, or post-consumer PLA materials can be recycled by chemical means to produce lactic acid monomers and oligomers. These materials can then be fed to the front end of a manufacturing process for making PLA lactide, ethyl
lactate, or other lactide derivatives. The recycling can be carried out with water at a wide range of temperatures (100–250°C). The reaction rate is enhanced by a catalyst such as nitric acid as is common in the PET recycling industry. The reactor residence times for PLA hydrolysis are in the order of hours, and depend on reactor temperature and catalyst level.

Chemical recycling should be considered in any waste management system for PLA, since from a life-cycle perspective, it represents a relatively small amount of net chemistry compared to the CO₂-to-PLA cycles for incineration or composting. Simple hydrolysis can turn waste PLA back into fully functional lactic acid, at potentially low economic and environmental cost, contributing to the total sustainability of PLA production.

The inferences from the studies of Vink et al.⁴³ confirm the positive impact that PLA has in addressing the key environmental concerns of today when compared to petrochemical-based polymers. They also show the additional benefits that can be gained by making environmentally responsible modifications to the existing manufacturing process. The commercial producer of PLA, Nature Works LLC, states that its commitment and ethos is based on ‘making plastics from annually renewable resources which meet the needs of today without compromising the earth’s ability to meet the needs of tomorrow’. Its philosophy is based on ‘reducing the environmental footprint’ and on ‘designing products with end-use disposal in mind’.⁴⁴ The company is committed to environmental responsibility and therefore it would seem that the polymer market as we know it today will experience some very significant changes in the near future.

5.6 Future trends

The vision for PLA fibers is more than just developing new performance products; it encompasses the additional goal of reducing total environmental impact. The result is a product that is more sustainable than comparable polymers on the market today. PLA fibers have made significant steps towards creating more sustainable products with unique performance attributes.

This technology allows one of the world’s most commonly used materials, plastics, to be made from simple plant sugars, which are then turned into fibers using conventional melt spinning equipment and processes. Made from annually renewable resources, PLA plastics and fibers use 20 to 50% less fossil fuel resources than is required by conventional petroleum-based resins. Fossil resource use in the manufacture of these materials will continue to decrease as plans are put into place to switch feed stocks to biomass (corn stalks, leaves, etc.) and possibly integrate alternative energy sources, such as windpower. With PLA, carbon dioxide is removed from the atmosphere when growing the feedstock crop and is
returned to the earth when PLA is composted. Since this process recycles the earth’s carbon, PLA emits less CO₂ compared to other petroleum-based fibers. Disposal of PLA fits within the existing disposal systems but also includes the additional option of composting.

PLA fibers are increasingly penetrating markets traditionally occupied by petroleum-based synthetics. The rationale for this trend is the increasing public concern over the depletion of natural resources and the accompanying atmospheric pollution. Various studies indicate that reserves of oil and gas will eventually be depleted, although the actual time scale is a topic of considerable debate. It is evident, however, that the use of annually renewable crops as a means of producing the materials we need to sustain our everyday lives will continue to accelerate.

PLA fibers offer the promise of ultimately reversing the damage we have imposed upon the earth while extending the usable life of the diminishing oil and natural gas reserves. Ultimately the technology of using natural crops to produce fibers and packaging will become widespread across the globe. The use of waste products, referred to as biomass, will also provide the fuel to drive the factories used to convert these polymers to the variety of products needed to sustain the ever-increasing demands for an improved quality of life. The key to this transformation lies in the economical manufacture of these renewable resource-based products coupled with meeting the performance demands of the targeted applications.

PLA fibers and packaging products are well on the way to realizing this goal. Presently these fibers are produced from natural corn sugar. However, as the technology develops, any starch-based crop will be used. Advances in farming practices will enable higher crop yields with subsequent improved economics. Additional improvements in the fermentation and polymerization processes along with economies of scale will also contribute to the improved cost basis for these products.

PLA products have come a long way from the early work of Carothers in 1932. The ability to economically manufacture packaging and fiber products has led to rapid growth. This growth will continue. We are at the beginning of a new industrial revolution in which PLA fibers are playing a leading role. In addition, PLA fibers are still in their infancy: improvements in chemistry and downstream process development will lead to new applications and a bright future for these renewable resource-based melt-spinnable fibers.

5.7 References

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Note
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6.1 Introduction

Every event has some impact on the environment. From the apocryphal fluttering of a butterfly’s wings in the Amazon rain forest to the detonation of a hydrogen bomb, a side-effect affecting our planet’s health occurs to cause an ecological imbalance that may or may not be recoverable. A balanced environment is essential if the Earth is to survive in a habitable form, and natural phenomena that have temporarily caused problems have been dealt with by the Earth’s natural functioning for millions of years.

Unfortunately, the advent of our technologically-skilled species has inflicted stress on the planet in ever-increasing proportions, with major consequences for its stability and survival prospects of all species on it, including humans. The textile industry, though not the only or most significant contributor, is exerting some impact and the contribution of polyesters and polyamides must be taken into account. Any aid that can be provided to reduce environmental harm will be of help in some small way to extending the life of the Earth.

6.2 Types of environmental impact

Environmental impact can take various forms (Slater, in preparation), some familiar and others not generally recognised. The former include energy consumption and pollution caused, together with global warming, melting ice-caps, rising sea levels and increasing frequency of adverse weather conditions. Less well-known problems include depletion of vital resources, rarity factors and ancillary effects like noise emission, radioactivity, odours, etc. There is usually little or no consideration of total impact, because most workers dealing with environmental concerns concentrate only on their own limited area.
6.3 Pollution types

Types of pollution that need to be recognised are encountered during production and use of polyesters and polyamides. The first is air pollution, in which harmful agents, including gaseous oxides of carbon, sulphur or nitrogen and particulate solids, such as dust, substances introduced during processing, or fragments of fibrous and other foreign matter, are released into the atmosphere. The gases may cause global warming, acid rain, toxic clouds, breathing difficulties, or other undesirable effects, while the particulate matter may exacerbate lung diseases, allergic reactions, visual obscurity, or soiling of fabrics.

Water pollution, the second possibility, occurs when salts, toxic agents, heat, chemicals, radioactivity, fibrous wastes or oil and petrol run-off are discarded. The chemical agents may affect food supplies or impinge on the natural world, while the heat and radioactivity can cause changes in life forms encountered. Solid waste can interfere with natural processes, such as stream flow and plant or animal access to sunlight, while oil-based materials can be absorbed into plants and cause diseases (such as cancer) in animals (including humans, directly or when animals are eaten) consuming the plants.

Land pollution arises when waste materials are discarded on to the earth’s surface. The materials may be harmful (chemicals or salts toxic to animals or plants, or causing a slippery surface) or may be harmless in this respect but damaging by occupying space and producing unpleasant sights or interfering with natural habitats of animals. Discarded substances can attract undesirable life forms (vermin, mould, bacteria, etc.) to bring about potentially harmful risks for human beings.

Noise pollution arises when the area around a site is subjected to loud sound levels. There are tables (Peterson and Gross, undated) of acceptable sound pressure levels that can be tolerated by human beings and above which exposure can cause deafness or psychological side-effects, but the disturbance of animals in their natural habitats is also a drawback.

Visual pollution is seldom recognised. Discarded waste materials fall into this class. It can also arise from permitting the use of billboards, hoardings or garish lighting in advertising. Coloured substances not in harmony with the natural world are significantly noticeable and hence pollution. In considering the polymers under review in this book, there are approaches that will reduce (but not eliminate) harm to the planet.

6.4 Pollution prevention and control

There are differences between prevention and control of pollution. Prevention implies that pollution is never generated, while control indicates that
pollution is inevitable but attempts are being made to restrict its quantity or effects.

6.4.1 Prevention
The only effective means of prevention is elimination of its source, by modifying production or re-using waste. Recycling, touted as a means of prevention, needs energy and produces waste. This is typical of human claims of pollution control; the pollution is not reduced or controlled, but merely hidden or transformed into another pollution, merely removing it from the vicinity where ‘reduction’ is taking place.

Waste minimisation can best be achieved by attention to detail in production. Enhancement of machine efficiency to avoid excessive friction, for example, can reduce pollution at the point of generation, or near the machinery if noxious fumes are also emitted. In addition, controls of temperature, humidity, quantities of reagents and other variables can lower waste. In re-use, reprocessing of ‘waste’ fibres is common, either yielding new textile goods or manufacturing a different product, such as artificial wood or road beds, that can use up the material and place it in a stable, non-polluting form into the environment.

6.4.2 Control
The attempt to control pollution (usually without significant success) is occupying much time in modern life. The aim is the subject of legislation in various regions, but the will to tackle problems of defining the situation is geographically varied. There is a lack of commitment in governments to do anything but pay lip service to the environmental lobby. There are plenty of examples, the most obvious being the refusal of several governments to ratify the Kyoto Accord. The earlier Earth Summit of Rio de Janeiro has also not produced any tangible action.

6.4.3 Legislation
Some legislation to reduce planetary harm exists. Germany has established regulations (Anon., 1998a) to limit or eliminate environmental and health hazards for textile and leather processing industries. Arias (1999) describes criteria that a manufacturer must meet to use the European Union’s eco-label, aimed at reducing air and water pollution and increasing human health and awareness of problems, as well as lowering water and energy consumption. McCarthy and Burdett (1998) provide details of labelling regulations for the industry and benefits of environmental awareness.

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It is interesting to examine work associated with the eco-labelling movement. Burdett (1997) notes that it stems from a system that was originally voluntary and widespread throughout Europe. Methods of testing were based on well-known standards, and additional national and ‘private’ systems were used. An anonymous author (1997) in India points out that, when firms compulsorily start adopting such practices, the standard soon becomes the norm. This can be expected to have a positive financial benefit for environmentally-based stocks, since products not meeting the new standards will be rejected due to the lack of eco-labelling. Morris and Crosby (1995) note, in an article on quality requirements for textiles and clothing in Europe, that legislative actions have taken place to improve public information on activity and the textile industry. These include laws on eco-labelling, eco-auditing, waste management and recycling.

A further anonymous article (2000a) points out that the textile sector must adapt to the concept of ‘Green Productivity’ to have products competitively poised in the market through eco-labelling, which would form the collective practices for this GP. Joshi (2001) carries out a case study on implementation of ISO 14001 in a textile unit based in North India, while Nadiger (2001) indicates that a price must be paid for eco-labelling, including a ban on certain chemicals in processing and containment of pollution to satisfy different eco-labelling schemes. In general, though, the legislation relating to textile production is weak and inconclusive, none of it being specifically directed at polyesters or polyamides. The major reason for this reluctance may be economic, since laws to provide adequate protection could decimate or eliminate synthetic fibre production. What is needed is a clear view of what damage can be tolerated, followed by laws to enforce limits more strictly.

Current regulations are unsatisfactory, because they are neither effective nor severe enough. It is true that there are laws about what can and cannot be done in releasing compounds, or engaging in activities with ecologically-undesirable effects, to the environment. These usually set limits for each substance or activity, regarding how much can be tolerated in a specific time. They include emission of air or water pollutants, and display of advertisements, signs or other types of visual pollution, as well as specifying the intensity of sound levels. All are excellent in intention but five drawbacks make them ineffective. These are non-uniformity of standards, lack of enforcement, insignificant penalties for flouting the law, disagreement on what is pollution, and lack of quantitative measures applicable to violation assessments.

The first arises from the fact that different standards are adopted in different areas. It is not too long ago that North American textile producers...
were shipping goods to Mexico or South-east Asia, where laws were lax, to have finishing carried out there that would have been illegal in their own countries. The finished goods were then returned to the original manufacturer to be sold as ‘domestically-produced’ ones. Some authorities are trying to improve the situation; this example is no longer applicable, because changes in laws in the former ‘dumping’ countries make it not worth while to pay transportation costs, even though standards there are still lower than in the host countries. Legislators are slowly responding to pressure from constituents to enact laws enhancing ecological responsibility, but changes tend to be ‘too little, too late’.

The second problem is lack of enforcement of standards. There are underlying reasons for this failure to protect the planet. First, there may be too few inspectors, usually as a result of shortage of money, to carry out effective policing. Second, the manufacturer may be part of a political lobby that threatens withdrawal of support if a government is too diligent in providing ecological protection, or there may be ‘financial encouragement’ in operation that convinces inspectors to turn a blind eye to contraventions of the law. There may also be conflicts of interest, where the manufacturer’s activities are profitable for legislators, who have no wish to curb harmful proceedings and persuade colleagues to agree with their lenient judgement.

Even where enforcement is evident, there is another reason why planetary damage continues. In many cases, the penalty applied for flouting pollution laws is relatively minor. If a manufacturer knows that, if convicted, he will only suffer a small financial penalty, he will continue to pollute and pay the insignificant fine regularly. If conviction frequency is low, he can write off fines as a business expense to pass on to his customers.

A further problem is the lack of agreement on what should be classed as illegal pollution. In most western nations, heavy emission from vehicles or factories is unacceptable, and the owner of the source is fined or forced to cease operation. At the same time, authorities in many countries of Eastern Europe, Africa, South America or Asia disregard these sources of harm, and some cities there are shrouded in toxic fumes that would close down the municipality in less tolerant countries. The excuse offered is that such nations are still catching up with Western competitors, and can’t afford to attack the problem until their economies are stronger.

An important reason why environmental efforts may not be effective is that inability to measure pollution accurately may lead to lenient sentences. In most countries, innocence is assumed until guilt is proved. If legislation depends on a quantitative assessment of pollution that cannot be measured precisely, there is no point in prosecuting an offender unless the pollution is so excessive that there can be no possibility of mistaken measurement.
as a defence. Pollution that exceeds considerably the amount permitted may be overlooked, rather than risking losing a case.

6.4.4 Environmental auditing

To minimise difficulties of pollution assessment, environmental auditing has been developed. Its major benefit is that it enables unexpected changes to be detected and can pinpoint where pollution is produced or undesirable additives taken up by the textile goods. Investigation can detect the source of the problem, which can subsequently be corrected. There are, however, difficulties that arise.

In theory, the process measures (by weighing) every component that enters or leaves a process. Thus, if (say) six raw materials are used to make a product, the weight of all six materials entering is compared with the weight of product. There will, in general, be a difference in the two readings. If this is negative (i.e. weight of product is less than the sum of component weights), then unexplained loss of material has taken place. This could be from evolution of some by-product (an undetected gas, liquid or solid that has escaped the measurement process) resulting from burning of fabric, evaporation of water, differences in atmospheric conditions when measurements were taken, or fibrous material lost as fly. If there is a positive difference (i.e. product weight is greater than component weights), then unexplained gain has taken place. Possible reasons include (as before) a change in measurement ambient conditions, chemical reaction with air or moisture, or accumulation of foreign matter (such as dirt or an adsorbed chemical) on the fibres. Smith and Lee (1998) identify trace impurities that can cause pollution, such as fibres, metals, VOC, or toxic organics. The work is complex and one outcome is the evaluation of analytical methods for polyesters, polyamides and other fibres. Kalliala and Nousiainen (1999) develop an environmental index model based on life-cycle assessment to determine total environmental impact, including energy in laundering, of a range of fabrics. They find that, for instance, laundering 100% cotton needs 20% more energy than a 50/50 cotton/polyester blend.

The first practical drawback of environmental audits concerns accuracy. The technique demands very precise measurement to ensure that all materials are detected. If an operator fails to note the existence of a material, or fails to measure its presence accurately, the error indicates the presence of lost or acquired mass. Time can be wasted in seeking this non-existent component, adding to the cost of production. It is possible, too, for workers to falsify results in the belief that management representatives want to hear good news, or may blame workers for large quantities of pollution. The discrepancy may not come to light until an inspector checks the audit figures, finding that unreported defects are in fact present. He may then
levy a fine or other penalty that is more costly than fixing the problem in the first place. There may be situations where, no matter how good the audit, pollution cannot be rectified. If a vital process produces large quantities of undesirable by-products, society may be faced with the choice of accepting it or agreeing not to need the product. If the product is essential (as in military or space applications), then no amount of auditing or fines will solve the problem.

It is for situations of this kind that legislation to permit trading pollution quotas was developed. The principle is that a company producing less than its permitted quota can ‘sell’ its unused quota to a company with excessive pollution. The latter is allowed to exceed its own pollution quota by paying for the privilege of producing excess. This exempts the ‘dirty’ company from prosecution and enriches the ‘cleaner’ company, but does nothing to help planetary overload. The worst aspect is that, as more companies are developed, added pollution permits are issued, no effort being made to reduce the quotas issued to existing companies to prevent increases in overall pollution.

A further difficulty is the need for vigilance to ensure that new (or the same) problems do not crop up later. If detection of a problem leads to a solution that saves money for the manufacturer, results are welcome. If there is merely a need to remove pollution without financial benefit, the manufacturer is less willing to participate. There is, too, another aspect that needs to be considered; short life expectancy for a textile product can add to the pollution load on scrapping it.

### 6.5 Environmental impact of textile production processes and use conditions

McKenzie (1999) suggests that natural fibres are not necessarily less environmentally harmful than man-made ones, by comparing ecologically-relevant production steps for natural, regenerated and synthetic ones. In a later paper (McKenzie, 2000), she observes that, although demand for organic cotton has grown, consumers will not pay higher prices for environmental conservation. She notes (McKenzie, 1999) that ecofashion costs more until mass production output is attained, but there can be no mass production until demand rises, which will only happen when cost decreases.

The actual techniques used in production are described adequately elsewhere in this book, and it is not intended to repeat them in any detail in this chapter. Instead, attention will be paid to the environmental aspects of each portion of production to consider its overall contribution to the impact of the life cycle of the two types of fibre. However, there is evidence that the actual experience of working in a factory where polyester or
polyamide is produced might be harmful to health. Hours et al. (1986) study mortality rates of personnel in such a factory and find that exposure to phthalates, nickel catalysts and other chemicals can cause serious problems.

6.5.1 Fibre and yarn production

Both polymers are currently produced almost exclusively from oil, and the extraction of this resource is environmentally costly, requiring extensive drilling with high energy machinery and causing pollution in the form of discarded waste and damaging spills during shipping. Once the oil is gathered, it is separated into fractions by heat energy, followed by conversion of the appropriate fraction by chemical reaction (again using heat) into the polymeric starting material for fibre production. The operation uses large amounts of energy, produces waste gas, liquid or solid by-products and can be unpleasant for the people working in it. Thiemens and Trogler (1991) reveal that nylon manufacture releases nitrous oxide, a more harmful cause (by a factor of up to 200) of greenhouse warming than carbon dioxide. Zenker, Borden and Barlaz (2003) report that 1,4-dioxane, a human carcinogen, is formed as a by-product in polyester manufacture, suggesting the need for further research into the development of cost-effective biological treatment processes. Spinning and extrusion steps are responsible for ecological problems. Melt spinning uses complex (and hence environmentally costly to manufacture) machinery that consumes large amounts of energy. Extrusion can bring about degradation of the polymers if temperature control is inaccurate, so causing waste to be discarded.

6.5.2 Transportation

It is appropriate to consider a pervasive environmental hazard that is constantly repeated throughout textile production and use. Textile materials have to be transported by vehicles during and after production. Vehicles are environmentally costly, but they also have a more sinister effect. Like textile machinery, they are large and complex, with the customary resultant costs, but there is also fuel combustion to take into account. Exhaust gases contain toxic or carcinogenic emissions, and legislation has been introduced in various countries to ban some components (notably lead) by modifying fuel chemistry. Approximately half of all air pollution is caused by vehicle emissions (Ross, 1972) and, although the textile industry is not responsible for all these, it must bear a fair share.

Exhaust gases remaining after removing (if possible) dangerous products cannot yet be eliminated, so attempts are made to improve fuel
economy, but this is rendered ineffective by tremendous increases in the number of vehicles produced annually. The most critical emission gases, in terms of immediate damage, include compounds of sulphur, heavy metals and organic by-products of combustion, all environmentally undesirable for obvious reasons. Even if these could all be removed by some miracle of science, the unavoidable production of carbon dioxide, a greenhouse gas, would still make exhaust by-products a source of harm in the long term. Transportation, too, uses oil extensively, as a lubricant as well as a fuel source, and we have already seen how harmful a substance this can be.

Not only the shipping of goods, but also the transportation of people to and from work, has a contributory effect. Vehicles cannot travel without roads, and the establishment of these is a major cause of environmental difficulties, from the extraction of raw materials to the mechanical devices used to lay them down. Their presence also eliminates land on which plants for food (or textiles!) can be grown, or on which animals could graze, thus introducing a somewhat different category of environmental factor. Its contribution to the ecological cost of textile production should be borne in mind.

6.5.3 Noise

Two further problems, noise and dust, are common ones. Both cause difficulties for humans, as well as for the environment, but are different from factors so far considered, so should be given special recognition. Of the two, noise is more difficult to classify quantitatively. Noise can be defined as any sound which, because of volume, frequency, speed or harmonic content, produces discomfort or pain in the listener. The vagueness of this definition provides scope for ambiguity, but some sounds are undeniably and universally classed as noise. There are two criteria that should be used in deciding where the sound fits in the desirable/undesirable continuum, that of intensity (or volume) and that of ability to please. The two may occasionally be in conflict, so that a sound which should be classed as noise because it is painfully loud may occasionally be acceptable because it stirs some deep inner emotional chord.

Textile equipment can be included in such a category. If a machine is running smoothly, with a quiet and steady, gentle hum, then its sound might be regarded as pleasant by the engineer responsible for its maintenance. If volume is extremely high, though, to the point where it is painful to the hearing, or is a shrill, high-pitched note, pleasure disappears and noise is excessive, even to the most dedicated engineer. The test of volume and frequency can be applied to any textile machine to determine its position on the noise spectrum. As sound intensity rises, the scale of
measurement is arbitrarily defined (Peterson and Gross, undated) in such a way that an increase of 3 dB(A) represents a doubling of volume. The net result of this is that a sound intensity of about 30 dB(A) represents a comfortable conversational tone and one of about 140 dB(A) is loud enough to cause physical damage instantaneously in the ears of somebody exposed to it.

In most industrial countries, the danger of subjecting people to high intensity sound has been recognised, and legislation exists to limit the exposure time of workers. Typically, people are not legally allowed to remain in a noisy area at 90 dB(A) for more than eight hours in the working day, and may not be allowed to enter the area at all if sound intensity is 115 dB(A) or higher, with shorter times being permitted at intermediate levels. The legislation also permits exposure at higher levels if approved hearing protection is worn, though there are two problems with this loophole. The first is that brain damage can occur via skull bone transmission even if the ears are protected, which means that the protection must include a helmet with acoustic insulation; the usual hearing protection, earplugs, is useless in preventing harm, even though it meets the law. The second problem is that of the machismo image. Because it is regarded by many textile plant operatives as ‘soft’ to protect their hearing, they refuse to wear appropriate devices unless forced to do so by supervisors.

Noise costs money. The cost may be direct or indirect, but it is considerable and should be recognised as a genuine financial, as well as environmental, one. The direct cost arises as a result of some process taking place in a machine. This may be an explosion, as in internal combustion, or frictional contact between two surfaces in a gear train or other mechanism. Noise is energy and as such must be generated by a power source. The electricity (or other fuel) used in the process cannot then be harnessed to carry out work, so is wasted. Money is paid to generate electricity, so the cost for this waste is irretrievably lost from profits. As a final point, energy generation is environmentally costly, and again the waste cannot be reversed.

Indirect costs are more subtle. Work carried out in various industrial premises has brought to light the suspicion (Slater, 1975) that hearing damage from noise exposure can cause social harm. Workers may experience boredom and loneliness or become withdrawn from social contact as their deafness worsens. In some cases, they may blame the machines that have caused their deafness, and deliberately take revenge by harming these perceived sources of a disintegrating lifestyle. Neglect of maintenance, slow correction of operating faults, or even deliberate sabotage, may occur in senseless acts of vengeance. All these cost money, and can exert a cost on the environment in the form of wasted energy or raw materials and excess exhaust gases.
6.5.4 Dust

Dust consists of tiny solid particles produced when materials break into fragments from mechanical, chemical or biological action. Because of their small size, they are easily blown about by air currents, or may float in the air for long periods of time without settling. In textile production, they are of two types, extraneous matter brought into the factory (as packaging, for instance) and fragments of fibres produced during processing.

Once they are released into the air, their behaviour can cause problems, an obvious one being the formation of a contaminant film on nearby surfaces. This necessitates frequent cleaning, requiring extra labour. When particles fall on machinery, they can interfere with operation, by increasing frictional force or lubricant viscosity and cause a breakdown if not removed. They can make a product dirty, so that it has to be washed (with detergents or cleaning materials and increased energy consumption) or rejected, causing significant environmental loading, as well as increasing production cost.

The second dust problem is health hazard, because of the chemical composition or size of particles. Cotton dust causes brown lung disease, while asbestos dust is responsible for lung cancer and dust from other fibres can cause allergic reactions, sneezing attacks, asthma and related diseases. Dust can interfere with plant health by coating leaf surfaces, preventing them from absorbing carbon dioxide and releasing oxygen. Polyester and polyamide production is not a major cause of dust hazards, but some effect is noted during later production steps (drawing, blending and spinning, for instance). For all these reasons, limits of permissible dust production are being established in the industrial world and applied to textile processes. Chellamani and Chattopadhyay (1998) stress the need to reduce fly and fluff production for workers’ health, and suggest control measures such as moisture addition, infrared lamps at strategic points, floating condensers, acid treatment of roller cots and use of overhead cleaners. Van Nimmen and van Langenhoven (1999), noting a sharp increase in contamination of fibres by foreign matter, such as plastics or wrongly-coloured stray fibres, give an overview of proposed solutions to the problem, with benefits and drawbacks of each one.

6.5.5 Fabric production

Once a yarn has been produced, there are several ways in which it can be made into a fabric. For polyesters and polyamides, these include weaving, knitting, non-wovens, fibre-to-fabric, film fibrillation, coating, laminating (Fung, 2002), needle-punching, bonding, tufting, and stitch-knitting. In every case, mechanical action creates risks of dust production, fibre
breakage, and hence waste. In environmental effects, familiar factors of machine size and complexity, and of energy consumption, still exist. The need for lubricants to reduce breakage (and hence waste) is also essential in today’s high-speed machines. In traditional looms, waste energy and loud noise are environmentally harmful, sound pressure levels reaching 110 to 125 dB(A), making the weaving shed dangerous (Slater, 1974) for hearing preservation of workers. Sizing to prevent yarn breakage and desizing also brings about ecological damage. Stöhr (2002) notes that an ideal size would be one that could be reused repeatedly, indefinitely and without limitation, and deliver optimum sizing results. He recommends a new material, UCF-4, which meets these criteria. Non-wovens, fibre-to-fabric, coating, laminating and bonding, using polymeric materials, hot glue or high pressure, also incorporate the same type of chemical loading, though an anonymous author (2001) describes a spunbonding and spun-lacing process using a single-step water-jet method to reduce its impact. Despite this claim, the usual hidden environmental costs already noted are inevitably present.

Tufting is one of the most widely-used manufacturing steps in which polyester and polyamide are involved, in view of the fact that most carpets are currently made by it. Environmentally, apart from the usual concerns about machine size and energy consumption, its most significant drawback is the need to apply glues or resins. These are not only toxic or otherwise harmful, but their subsequent treatment by heating to cure them can bring about gas evolution to create air pollution. Membranes, providing a waterproof and breathable fabric (Bajaj, 2001), involve sandwiching microporous polytetrafluoroethylene (PTFE) between two layers of fabric, usually polyester, to prevent delamination of the fragile PTFE. Making the fabrics and the adhesive needed imposes environmental costs. Coatings (Kubin, 2001) provide a compromise between microporous membranes and finishes and are felt to be more satisfactory (Fung, 2002) from the ecological perspective.

6.5.6 Fabric chemical processing

Once a clean fabric is available, finishing can begin. The type selected depends on the fibre content or end-use of the fabric, and synthetic fibres need fewer finishing treatments than natural ones. They do not, for instance, generally require protection against microbiological hazards or fire and, in other treatments, such as moiré finishing, machinery and energy factors (plus steam when it is used) are the only environmental costs. Protection from ultraviolet degradation is often needed and includes chemical substances (including amines, sulphonated or benzoyl
compounds and other complex organic reagents) that may be damaging to the planet.

The main cause of environmental concern in chemical processing, though, is coloration by dyeing or printing, and Christie (2007) has recently edited a volume dealing extensively with this subject. Benisek (1999a) provides a summary of eco-friendly dyes, but dyestuffs are frequently toxic or carcinogenic and, when released in waste waters from a plant, are visible. Natural dyes are ineffective for synthetics, so new synthetic dyeing techniques with lower planetary loadings are being developed. Swett, Thwigg and McCaleb (1984) compile a data base of 1434 commercially-significant dyes and pigments used in the USA, providing information on toxicity, economic costs and producers. The use of disperse dyes, plasma treatments (Wakida et al., 1998; Anon., 1998b) or supercritical dyeing (Lennox-Kerr, 2000; Kawahara et al., 2001) is recommended by various other workers to reduce economic and environmental costs. Anis and Eren (2001) note that the use of reactive dyes for improved washfastness carries an unexpected environmental impact of added rinsing needed to remove excess dye not absorbed by fibres. They report the development of new bioreactive molecules to reduce (but not eliminate) this drawback. The same authors (Eren and Anis, 2005) later compare the environmental impact of polyester alkaline dyeing with five different after-treatments, noting that the technique is successful in reducing harmful effects if the correct dye and after-treatment combination is used. Paylor, Ayers and Bell (1986) propose biological techniques in laboratory and field trials to demonstrate that chemical testing alone is unreliable in predicting toxicity level in a mixture of textile and domestic waste. Kwok, Xin and Sin (2003) attempt to predict toxicity of dye effluent and create a toxicity index to aid in the aim of reducing environmental damage. Riva and Valles (1994) attempt to determine the toxic effects of surfactants used in the mothproofing process on freshwater microalgae species, since no toxicity data exist for these compounds. Bae, Freeman and Kim (2006) examine the influence of new azo dyes on the aquatic system, showing that the copper incorporated in them has an adverse impact and suggesting a new method to evaluate the effects of dyes on the aquatic environment.

Printing uses the same reagents as do dyeing treatments, together with a printing paste that can cause water pollution. In compensation is the fact that a printing paste can be more easily collected, with lower amounts lost, than can a dye liquor. In addition, because the purpose of printing is to force colours into fibres, there is less need to wash out excess colour, so leading to lower quantities of discarded pollutants. Pollution from coloration occurs because, when dye is applied, it is not all picked up by the fibres and there are inevitably residual amounts that cannot be adsorbed.
Despite efforts to recycle them, large quantities cannot be re-used, either because the shade is not applicable for the next fabric batch, or because dilution is too great to make recovery economically viable. Flock printing and heat transfer printing overcome this drawback, but ecological aspects of fibre dyeing and fixing agents have to be considered in the former case, while the need to make special paper and a separate printing stage using heat for dye application, together with disposal of the paper, are important in the second. Both methods, though, remove the need for liquids and printing pastes, and omit washing, steaming and drying steps, so reducing ecological harm.

Emissions to water are the obvious example of substances released into the environment, and a report (Anon., 2000b) confirms that waste water discharges from wet processing are toxic and likely to have immediate or long-term harmful environmental effects, though ‘smart rinsing’ (Bradbury et al., 2000) can provide more effective use of waste. Reife and Freeman (2000) summarise possibilities of pollution prevention by waste minimisation and source reduction in producing dyes or pigments, while purification before discharge is attempted (Bischefberger et al., 1999; Canziani and Bonono, 1998; Ibrahim et al., 1998; Papić, Koprivanoc and Božić, 2000; Gonçalves et al., 2000; Slater and Barclay, 1998), using sand filters, cross-linked wood sawdust composites, coagulation/flocculation processes, anaerobic sludge blanket reactors or inorganic clays with other materials to absorb dyestuffs.

Air emissions should be mentioned briefly. These can arise from many segments of the industry, and their removal is often attempted before discharge to air or water. Recovery of waste process heat (anon, 1999a) and air cleaning installations (Anon., 2002; Freiberg, 1998) can help in reducing environmental impact.

A contemporary preference in many cases is to try to minimise resource depletion and pollution production as much as possible by re-using or recycling materials, these being extended (Benisek, 1999b) beyond water treatment to all materials used by the industry and including biological treatment of waste water, digestion techniques, neutralisation of alkaline effluents, reduction of mothproofing effluents, spray dyeing and fibre identification. The recycling of fibre or fabric materials is also regarded as useful (Kohler et al., 1999; Raje and Rekha, 1998; Diounn and Apodaca, 1999; Methner-Opel, 1998; Roberts, 1999), with interest ranging from fibres and fabrics to end-products. Polyester figures in some of this work (Mannhart, 1998; Hansler, 1999; Goynes, 2000; anon, 1999b) and carpet polyamide recycling is also prominent in publications (Anon., 1998c; Realff et al., 1999; Tullo, 2000). Woolridge et al. (2006) show that, for every kilogram of recycled polyester used in garment manufacture, energy consumption is 90kWh lower than when virgin fibre is used.
6.6 Use conditions

Even after the textile end-product has been manufactured, it may be a cause for concern regarding its ability to cause environmental problems. When a product is discarded, it can be a source of harm to the environment. First, it is a visual pollutant if thrown on to the earth’s surface. If it is burnt or buried, it can inflict damage, as a source of toxic or greenhouse gases in the first case or harmful decomposition products in the second. Polyesters and polyamides are particularly damaging in this respect, since they tend not to be completely biodegradable. Long life expectancy is thus a major factor in reducing the environmental impact of textile products.

6.6.1 Apparel and household goods

The most common reasons for discarding apparel relate to consumer dissatisfaction and there are various reasons for this. Comfort, whether physical or psychological, is a common cause for complaint and synthetic fibres are often regarded as problematic. Perspiration discomfort in high humidities, or static shocks at low ones, are cited as drawbacks unless adequate finishes have been applied. Even then, comfort level can decrease gradually if the finish is removed by continual laundering. In compensation, synthetic fibres are usually more durable than natural ones, so will not be discarded as quickly if comfort problems can be solved. The same is true for household goods, which are not usually as critically affected by discomfort problems due to humidity. Sources of degradation responsible for reducing life expectancy include heat, light, and chemicals used in cleaning or maintaining fabrics; again, polyesters and polyamides have advantages over natural fibres as they are less easily harmed.

6.6.2 Industrial textiles

Industrial textiles tend to be used in more severe situations than domestic ones. They are generally required to have better resistance to weathering and mechanical attack, because they are normally used in outdoor conditions with abrasive, tensile or other stresses imposed on them. Again, synthetic fibres are preferred for superior strength and resistance to biological degradation. Construction to give a stronger fabric and finishing to impart resistance to ultraviolet degradation are arguably the most critical factors in enhancing durability.

Sollinger, Levsen and Wünsch (1993) estimate indoor air pollution by emissions of volatile organic compounds (VOCs) from polyamide textile floor coverings, while Kriek et al. (2001) study VOC emissions taking place
during polyamide resin production. They calculate an emission factor for each set of processing conditions, to be applied by a scaling operation from laboratory results to commercial manufacturing situations as a predictor of the quantity of each substance released as a fraction of the resin produced.

### 6.7 Pollution control strategies

Current attempts to control pollution are aimed in general at a few areas of concern, these being primarily air, water and (to a lesser extent) land or noise problems. The usual technique is to establish permissible limits, then try to enforce legislation that ensures these are not surpassed. As mentioned earlier, this approach is unlikely to be successful, so others must be tried.

#### 6.7.1 Air pollution

Rey (1998) provides guidelines on the technology available for controlling smoke and odours from textile finishing, and Holme (1998) recommends a three-faceted method (considering emissions, efficiency and economy) for pollution reduction or prevention. The main objection to these lines of approach, their uneven application in specific regions, is still present. Smoke stacks continue to pour out noxious fumes, but the traditional solution, still tolerated in some countries, has been to build the stack higher to allow toxic gases to be dissipated further into the atmosphere. Legislation tends to accept this compromise, yet it is not effective for lowering pollution, as is becoming more and more obvious as the number of stacks, and the amount of gas discharged from them, increase steadily. Air pollution is not localised; harmful gases are carried by air currents to other regions and, eventually, all parts of the world. Absorption is no solution, since it merely transforms the air pollution into another type when the neutralised gases are discarded on the land.

There are other examples of global dissipation of environmentally-dangerous substances. Dioxins and furans, among the most toxic chemicals known because they can bring about poisoning or adverse genetic mutations in extremely small concentrations, can be emitted from textile finishing treatments. Their discharge is, in theory, strictly forbidden in most developed countries, yet there are frequent reports in these very countries of their appearance in waste streams; it is cheaper to discard them, as long as their source is not detected, than to deal with them effectively and responsibly, simply because their destruction is technically difficult and hence costly.

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6.7.2 Waste water treatment

It is in waste water treatment where there is greatest concern, and hence where the greatest amount of interest is focused. Water treatments range from simple to complex, on continuous or batch basis. The simplest way of dealing with the problem is to avoid creating it, and some workers have suggested modification of production to lower environmental impact. Anis and Eren (2003) use an oxidative clearing method with mononicotinotetrazine reactive dyes to eliminate fastness problems when a one-step polyester/cotton dyeing process is used to reduce the amount of water discharged. The same authors (2004) later discuss the selection of suitable dyes for this process and specify rigorous conditions that have to be met. Fan, Hoskote and Hou (2004) note that batch dyeing of polyester and nylon usually involves the discharge of large quantities of waste dyestuff and suggest lowering effluent by optimising pH and salt control or by re-use of spent dye. Yin and Jiao (2005) adopt a new sizing mixture for polyester/cotton blends to reduce cost and environmental damage.

Supercritical dyeing, already mentioned, is commended for reducing ecological impact. Draper, Montero, Smith and Beck (2000) measure the solubility of various disperse dyes in supercritical carbon dioxide, while Montero, Smith, Hendrix and Butcher (2000) provide an overview of the process. They note that, if carbon dioxide can supplant water as dye carrier, there is no liquid effluent to discard (though, of course, there is an escape of some greenhouse gas). Hendrix (2001) expands on the idea, noting added benefits, such as the lack of need for any drying and hence the reduction in economic, as well as environmental, cost. Montero, Hinks and Hooker (2000) continue the work by providing a typical example of how cyclic trimer deposits can be reduced in a supercritical dyeing process.

Once discharge of a pollutant has occurred, steps are possible for reducing its impact. They can be classed in three stages of effectiveness. In the first, a simple mechanical filter is placed in the path of the water requiring treatment. This removes solid contaminants, the size of fragments removed being dependent on the mesh of the filter. Waste cloth, fibres, or solid detritus are examples of pollutants removed by this treatment.

The second type includes agents, such as clays, with surfaces on which pollutants are adsorbed. This is suitable for dissolved substances and particulate matter too fine for filtration to be applicable. Barclay (2000) surveys the available techniques and reports the use of sodium bentonite and activated carbon, together with flocculants, such as iron or aluminium salts, dissolved-air flotation, or organoclay derivatives. Continuous treatment techniques in this category tend to be less useful than batch ones over time, since the active agent in the treatment needs to be replenished occasionally and its effectiveness is reduced as its life cycle progresses. Batch
processing, though, is more expensive, because it is slower and uses more removal agent than is necessary to ensure that all pollution has gone. In terms of effectiveness, Barclay’s work indicates that removal of dyestuffs, mordants, suspended solids, silicone emulsions and resin finishes can be almost complete, depending on the treatment regime chosen, and that there are specific optimum removal processes for each type of pollutant. Uner et al. (2006) evaluate a coagulation-flocculation on a COD-based molecular size for a textile finishing mill effluent in an effort to establish optimum treatment alternatives.

In the third approach, a chemical agent is added to the water to bring about some kind of physicochemical activity that removes residual pollutants by precipitation or chemical reaction, such as oxidation. This is the basis of water treatment at the municipal level, but can be used if necessary in the plant before an effluent is discharged.

The next technique to be investigated recently is the use of light radiation as a mechanism for degrading dyestuffs. Chaturvedi et al. (2003) use three types of reactor and a continuously-stirred batch operation in sunlight, finding reductions of up to 99% in colour and 84% in COD under the optimum combination of dye and photocatalysis conditions. Arslan-Alaton and Dogruel (2004) combine photodegradation of dye effluents with silicadodecatungstate as catalyst, finding that regeneration of this catalyst (a major stumbling block in earlier techniques) is simple. The same two authors, with colleagues, also use (Uner et al. 2003) ozonation to treat dyehouse effluents, a process evaluated by Radetski et al. (2002) and found to be relatively effective in reducing toxicity.

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out a quantitative evaluation of a bioreactor to allow re-use of textile effluents to be considered. In a review of bioreactor studies, van der Zee and Villaverde (2005) discuss the effectiveness of a combined anaerobic-aerobic treatment technique for use with azo dyes.

All these methods of treating effluent water are accompanied by environmental consequences. In every case, the treatment agent must be manufactured, by mechanical processing or by chemical production techniques requiring machinery, raw materials and energy. Chemical treatment may bring about unanticipated reactions, the usual example quoted being the production of dioxins or other toxic substances when excess chlorine is used to treat drinking water. In all cases, the waste removed, in the form of solids discarded to the land or chemical agents that need disposal, cannot be ignored. Their removal is again an example of the fact that pollution is never ‘removed’ but merely transformed into another type or transferred to another location.

6.7.3 Land pollution removal

Land pollution falls into two types, visible or unseen. Visible pollution includes solid waste, thrown on the ground deliberately or blown there by winds, and waste located at specific landfill dumps. In addition, waste products dissolved in water can be discarded on the land and soaked up by it, thus being hidden (except in the case of some coloured agents staining the ground) from direct observation. In general, the latter type is more dangerous than the former, since its presence is not recognised, so cannot be counteracted, and tends to include harmful substances easily transferred to underground sources of drinking water.

Removal of visible waste can only be achieved by collection, either by hand or (if the unit size is large enough) by a mechanical device like a suction vehicle or rake. Hand removal is not often used because of financial cost, but mechanical techniques are dependent on machinery manufactured by environmentally-costly processes and using energy during both manufacture and use. Hidden waste is much more difficult to remove. If it is dissipated into the water table, it is impossible to retrieve it and water has to be treated before using to avoid poisoning by oral ingestion or skin contact absorption. If the pollution is localised (as an oil spill, for example), there is a possibility of removing it. In large-scale spills, the usual approach is to break up the oil by means of detergents, but this is just another instance of hiding pollution, since the small globules are still present even though invisible. For smaller spills, soil remediation is the customary solution, pumping a solvent into the ground then extracting the solution before separating the oil from the solvent. Again, this is expensive, financially as well as environmentally, so is less than satisfactory. Newer techniques use
supercritical carbon dioxide for extraction, an approach which is less costly but still leaves a greenhouse gas.

6.7.4 Noise pollution control

The best way to control noise pollution is not to produce it in the first place and it makes economic, as well as environmental, sense to prevent its initial production. Noise is created when gases expand rapidly or when surfaces are rubbed together under frictional contact. Control of noise can therefore be attempted by supplying expansion chambers for releasing gases at a lower rate, or by lubricating contacting surfaces. Expansion chambers must be designed for each application, and the viscosity of lubricants must be selected carefully to ensure that high frictional contact between the surfaces is prevented without hindering machinery operation by having too viscous a separation medium. In textile processing, new techniques using shuttleless looms reduce high noise levels in weaving.

6.8 Eco-friendly technology options

The term eco-friendly has been coined to define a process that is effective without harming the environment, an impossible aim since all processing is damaging to some extent to the earth. McDonough and Braungart (2002) claim that a ‘not too bad’ solution to an environmental problem is worse than a ‘bad’ one, since the former lulls us into a sense of false security in which we pay lip service to environmental responsibility without achieving any satisfactory goal. They note that the common trust in the effectiveness of reduction, recycling and reuse is misplaced, since each tends to use added energy (even if only for transportation) and more resources to achieve a new end-product that is inferior in properties in comparison with the starting material and has toxic or otherwise harmful reagents added in the transition from original to new material.

Thus, the permissible limits set by various authorities on air emissions (and their control) are not a solution to the pollution problem. All that is achieved by lowering emission levels is a reduction in the rate of planetary destruction. We have reached the point where the earth’s corrective mechanisms are incapable of dealing with the load placed on the ecosystem and we are compounding the problem by removing aids to correction (such as trees or soil particles) that are the only means of reducing pollution by natural processes.

The subject of eco-friendly technology, as it relates to polymers, is reviewed by Khare and Deshmukh (2006) in a report on attempts to reduce the impact of plastics on the environment. They include substitution of non-renewable resources, such as petroleum, by bio-materials, but ignore
the inevitable (and large) environmental costs of producing these latter. Bajaj (2001) focuses on the challenges facing the textile finishing sector in an overview of various eco-friendly treatments. The same author later (2002) reviews changes that have taken place in finishes designed to meet consumer demand in such areas as comfort, ease of care, health or hygiene without causing as much environmental damage as the traditional techniques. Ibrahim et al. (2005) discuss chemical pre-treatments that they feel can help to maintain eco-friendly pigment printing with enhanced colour depth. Shin, Son and Yoo (2007) show that plasma grafting can be used to increase dye uptake, as well as moisture regain, for polyester, thus possibly providing potential reduction in ecological impact by reducing the treatment time and types needed. At the same time, of course, the comfort aspect is improved, serving to prolong life and hence reduce environmental impact.

6.8.1 Biodegradable polyesters and polyamides

Natural fibres, because they are degraded quite rapidly without producing harmful by-products, have long been considered more eco-friendly, and much work at present is focused on the effort to manufacture polymers that achieve the same type of property. Unfortunately, easy degradation goes hand-in-hand with short life, defeating one of the major benefits of polyesters and polyamides. In addition, emissions to water during the production of five types of biodegradable polymers are shown by Arfsten et al. (2004) to inhibit plant growth.

Problems arise because the so-called biodegradable synthetic fibres at present are not totally and easily biodegradable. The traditional technique is to produce fibre molecules composed of an admixture of polymer with starch particles that allow breakdown of the grain structure to take place rapidly. This ‘solution’ is highly undesirable. When the fibre is discarded, weathering or other degradative mechanisms destroy the starch, but the polymer is still present. The microscopic particles, though invisible, are so small that they present a large surface area to the environment, making them susceptible to rapid decomposition. Thus, the toxic end-products of decay are released far more quickly and are able to contaminate their surroundings at a much higher rate, making the situation more dangerous than it would otherwise have been.

Recent efforts are mainly directed in three avenues of approach, photodegradation, bacterial activity and the use of fungi. Andreopoulos and Theophanides (1994) compare the suitability of photodegradation and biodegradation technologies for speciality applications, reviewing research on degradable polymers at the same time. Andreoni, Baggi, Guaita and Manfrin (1993) note that three mixed cultures of aerobic bacteria are able
to grow on low-MW polyamides, but a MW above 11 000 inhibited growth. Growth was greater as MW decreased and appeared to use cyclic and linear oligomers of up to eight monomeric units present in the polymeric matrix. The work on photo-oxidation does not appear to have been pursued beyond that preliminary reference yet, and bacterial degradation seems to be popular at present.

Matsumoto and Doi (2003) suggest that the best way of dealing with the problem is to use a family of polyesters actually synthesised originally by bacteria, which are thus readily biodegradable. Their main drawbacks are, first, their high cost and, second, their inferior properties in comparison with petrochemical-based polyesters, and the authors are carrying out research to attack both of these problems.

Zheng and two colleagues (2005) review the topic of plastic waste biodegradation and note that aliphatic polyesters are easily attacked by microorganisms directly because of the potential hydrolytic cleavage of ester or urethane bonds in their structures, in contrast to aromatic polyesters. Gu (2003) feels that, although our understanding of polymer degradation has advanced recently, the subject is still inadequately addressed, as indicated by the lack of information on mechanisms and microorganisms involved. New techniques and tests currently emerging are also described briefly. Teeraphatpornchai et al. (2003) screen microorganisms isolated from soil samples for their ability to degrade various polyester-based plastics. The most active strain, designated as TB-13, was selected as best, but requires additional carbon sources in the form of enzymes to thrive. Kim and Rhee (2003) report the development of a variety of biodegradable polyesters to reduce the environmental impact of waste plastics. They confirm that aliphatic polyesters are more easily attacked than aromatic ones, but suggest that fungi, rather than bacteria, may be more likely to attack recalcitrant molecules. They note, though, that work on these agents is scarcer than that on bacterial degradation and report recent advances in knowledge.

6.8.2 Water conservation techniques

One of the more useful ‘earth-saving’ processes is reducing water use, and the textile industry takes an active interest in developing this form of environmental economy. Heat transfer printing, already mentioned, is a case in point; the costs are not so severe that they outweigh the reduction in damage achieved by removing the need for using water and releasing dye solutions. A second positive step is the substitution of supercritical dyeing, also mentioned, for aqueous methods. As long as the pressure can be maintained and the major part of the carbon dioxide (or other solvent) can be recovered, the ability to achieve deep and even shades of dyeing without water (and without releasing aqueous dye solutions into the environment)
is an excellent step in the right direction. Knittel and Schollmeyer (1995) note, when recommending supercritical fluid dyeing of polyester or polyamide to eliminate water use, that the technique costs less, avoids the need for auxiliary agents and saves time.

Environmentally-friendly dyeing has also been an aim of the industry for a number of years. Tajiri and Matsui (1973) conduct laboratory and pilot plant experiments to study the recovery of lactam monomer from waste water in nylon manufacture, using activated carbon. Schoeberl and co-workers (2005) take the further step of recycling the textile waste water itself, using ultrafiltration instead of the more complex and expensive bioreactor membrane. They suggest that the idea has the potential to cut down water use by 87.5% in washing, that COD is reduced by 80% and that consumption of washing agents can be lowered by 20%.

A further advance is the result of increased care taken over a process. It is possible, by good process design, to minimise costs by using the correct amounts of chemical agent and energy to bring about the desired reaction. Measurement precision is easily attained with modern equipment and can be used, in conjunction with minimal processing temperatures, to achieve significant reduction in planetary harm. A further suggestion for planetary conservation, still experimental, is the idea of making polyester or polyamide from a natural source, corn being the usual example quoted. Since corn is a renewable source, the argument goes, fibres can be made from it without harming the earth. Sadly, this is not true. The growing and harvesting of corn for food already puts a tremendous strain on the planet, because of the fertilisers, herbicides, pesticides and equipment essential by modern standards, and subsequent processing to separate the husks and convert them into fibres can only be done at considerable cost to the eco-system. All the process actually achieves is a reduction in the use of oil and, although this is a laudable aim, the drawbacks of the complex processing are an unacceptable price for this end-result. Yet another drawback is the fact that land potentially available for food growing is sacrificed to make fuel.

6.9 Future trends

Future prospects will be controlled by environmental concerns. We are at a stage where planetary damage is approaching the point of no return, one in which an uncontrollable runaway state is almost upon us. The passage of time merely serves to bring to light more examples of the way in which human damage is destroying the earth’s ability to cope with our presence, and there is a growing tide of opposition to actions failing to take into account environmental health. Thus, the textile industry of the future, if it is to survive, must strive to reduce ecological harm. Dixit (1998) feels some
companies may be forced to close because of inability to meet the ecol-
ogical challenge and emphasises the importance of safer and better treatments
for reducing pollution to satisfy newly-emerging regulations. Shaver (1999)
stresses the need for companies to enhance management of environmental
responsibilities and feels that this will be accompanied by a reduction in
costs.

For example, financial benefits of ecological responsibility, in the form
of tax deductions (Battersby, 1999), improved production (Moore, Money
and Orzada, 1999), cost savings from water recycling (Wakeling, 2001;
Hohn 1998) and benefits of new design (anon, 2000c) are reported. There
are, though, costs that cannot be ignored, financial as well as environmen-
tal. The energy taxes proposed in the EU (Russell, 2001) will be heavily
biased against the textile industry because its processes are energy-
intensive. Taxes also allow increased pollution to continue as long as the
producer (and hence the consumer) is prepared to pay a higher price for
the privilege.

6.10 Sources of further information and advice

A more detailed discussion of many of the points made in this chapter may
be found elsewhere (Slater, 2003), but some critical facts need to be
stressed. First, the emission of any harmful compounds, especially some
of the breakdown products mentioned, is undesirable, so the industry must
recognise that the earth cannot tolerate this abuse indefinitely. Because
emissions are similar in different regions of the earth, standards of safety,
or of environmental protection, need to be harmonised over the planet to
be effective. If there is a difference, then an unscrupulous manufacturer
(as mentioned) will merely move ‘dirty’ production activities to a part of
the world where standards are less severe to take advantage of lax laws
there, and will continue to pollute the planet rather than clean up produc-
tion methods. This activity must be forbidden if the problems of environ-
mental irresponsibility are to be overcome.

Second, one point seldom made is that the cure for an environmental
problem may be environmentally harmful. If, for instance, a specific
machine or reagent is needed to remove a pollutant, then the cost of manu-
facturing that machine or reagent provides a burden to the planet that is
usually ignored. The result of its use, too, may leave a pollutant of a dif-
ferent kind; absorbing an acid gas in a chimney stack by scrubbing through
a water trap, for instance, leaves an acidic solution to be discharged to the
water system. If an alkali is used to absorb the gases, or to neutralise this
solution, then a salt is produced, to be deposited on the land. Pollution is
again not removed, merely disguised to meet restrictive legislation. Thus,
because the total effect on the planet is undiminished, many efforts put
forth are ecologically useless. This is true at all stages in the manufacturing process; wherever a problem is ‘solved’ by any method other than reduction of consumption or production, the ‘solution’ leaves behind a residue that can create an ecological nightmare when disposal is attempted.

At any stage in a production train, there are factors that do not belong to the step being considered. All raw materials, for instance, are assumed to be present as if by instant creation, as are items of equipment needed to carry out the step. Nevertheless, each of these has an associated environmental cost that should be taken into account for complete evaluation of the step. The principle of the analytical procedure has been described in detail elsewhere (Slater, 1994) and leads to a complicated process of iterative calculation that can trace costs back to the initial step of extracting minerals from the earth for the production of metals or polymers used in manufacturing the equipment. Additionally, the costs of acquiring and using the sources of energy necessary to operate the machinery at all stages in production, from fibre and iron ore, etc., to working equipment and yarns, can be derived. The net result is that no process is free of environmental cost. If one is strictly pedantic, even the production of carbon dioxide (an important source of global warming) by people breathing can be considered an environmental cost. When the need to provide them with food, housing, transportation, working space, and other luxuries used at the planet’s expense are taken into account, it becomes more and more clear that human beings are a major source of environmental cost, whether or not they operate textile machinery. The crucial point about this method of analysis is that, if the production at any stage is faulty, the cost to the environment does not just include that of wasting the product of the step where it is discarded, but also the entire production train to that point. The later in the overall process a rejection occurs, the more cost to the environment is involved. This is a parallel cost to the financial one, fortunately, which leads manufacturers to avoid such waste by finding an alternative use for substandard materials instead of rejecting them.

6.11 References

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Part II

Improving functionality of polyesters and polyamides
Specialty fibers from polyesters and polyamides

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7.1 Introduction

Out of the many polymers developed for synthetic fibers, only a few dominate the commercial market. Polyesters and polyamides are the most widely used textile fibers. Polypropylene fiber has a major presence in nonwovens and to some extent in carpets and rugs. These fibers are commercially produced in large capacity plants, available in sufficient quantities, and are priced relatively low. Overall, these industries appear to have smooth sailing ahead since demand is increasing due to the growing population, but there are issues such as overcapacity, increased competition, and narrowing profit margin. Some manufacturers are diversifying into specialty markets by creating a series of new products using the same machinery with minor modifications in the process or equipment. These specialty fibers are focused on high value products that are new and improved over current materials, in both existing and new applications.

Typically, commodity fibers have a circular cross-section. Other cross-sections such as hollow, trilobal, hexagonal, etc., are considered specialty fibers. This modification can be accomplished by installing spinnerets with the desired capillary configuration. Due to the change in the cross-section, they exhibit amazing properties. For example, hollow fibers have better insulation properties, and trilobal cross-section fibers have a sparkle. Similarly, there has been growing interest in the specialty fiber markets comprised of two polymers, known as bicomponent fibers. These bicomponent fibers are produced by melting two polymers of differing structure separately, bringing the constituent polymer melts together in the die capillary, and fusing them into the forming fiber. A bicomponent fiber with a PET core and an HDPE sheath is a very popular binder fiber in nonwoven production.

Commodity fibers are generally made from homopolymers. The specialty fibers are produced from the polymer through chemical modification. Modifiers are added at some stage of the polymerization process or
during polymer extrusion. Cationic dye-able polyesters, flame retardant polyesters, and biodegradable polyesters are examples of modified polymers.

Specialty fibers are born out of research and development efforts tailored to meet the application-specific needs. This is accomplished if the modifications are easy to implement, and modifiers are easily available, exhibit better performance attributes, cover wider needs, and the products are readily available to the user. Special considerations at the design and manufacturing stages of these specialty fibers are discussed in this chapter.

7.2 Production, properties and applications of physically modified fibers

7.2.1 Bicomponent fibers

Bicomponent fibers are comprised of two different polymers of differing structure melted separately and spun together into one fiber. The constituent polymer melts are brought together in the die capillary and fused into the forming fiber. The design of the spinneret-pack assembly plays an important role in the quality and cost aspects of this product. This involves additional capital cost for the second polymer melt system and the special spinneret packs. Furthermore, cleaning, maintaining the parts, and inspection of these spinnerets are more expensive compared to monocomponent spinning. More common types of bicomponent fibers are (as shown in Figure 7.1): sheath-core, side-by-side, segment pie, and islands-in-the-sea

![Diagram of bicomponent fiber cross-sections](image_url)

7.1 Bicomponent fiber cross-sections.
bicomponent fibers. Advanced Fiber Engineering has developed nearly 400 specialty capillary designs for such applications. Advanced fabrication techniques are used in achieving capillaries to produce fibers with various shapes providing high shape factor or high specific surface. This increased surface area is useful in trapping dirt, with hollow cores of desired shape to enhance permeability and acoustic behavior, and perfect geometry to impart desired feel, stiffness, and light effects (luster, sparkle etc.).

Sheath and core

The largest commercial use of bicomponent fibers of the sheath-core type is for self-bonding binder fibers consisting of low melting temperature sheaths and higher melting temperature cores. Common sheath/core combinations in such applications include PE/PET, Co-PET/PET, and PP/PET. These fibers are usually blended with commodity fibers in the desired proportion to produce nonwoven webs by carding, air laying or any other suitable technique. Such webs can be through-air bonded or calendered. The sheath provides the fusing point while the core preserves the integrity of the fibers in the nonwoven. In these cases, the core always consists of a higher melting polymer than the sheath. The products containing these binder fibers can be processed at temperatures ranging between the melting temperatures of the sheath and the core. A major advantage is that it creates a nonwoven free of chemical bonding agents and is widely used in through-air thermal bonding of natural fiber or staple fiber webs.

A second type of sheath-core is filled fibers consisting of a core product produced from either cheaper material, or expensive materials (such as metal, dye) to impart desired special properties as well as esthetics to the final product. Improved conductivity is achieved by having a metal or carbon core in the bicomponent fibers. A third type of sheath-core is filled fibers consisting of a sheath product produced from expensive material (such as heat or chemical resistant polymers) to impart desired special properties as well as to keep the product economical.

In both the second and third type of bicomponent fibers discussed above, the desired property or functionality is on the surface and the additive does not have to be in the whole bulk of the fiber. Since specialty additives are generally quite expensive, reducing the amount of needed additives leads to considerable cost savings.

Side-by-side

Side-by-side products are typically used as self-bulking fibers. Self-bulking is imparted by two polymers having different shrinkage properties. These self-bulking fibers have a higher bulk and have found use in staple fiberfill
applications. Recently, they have been used in textured upholstery filaments. This technology is gaining popularity due to its higher processing speeds, and reduced cost. Sometimes the side-by-side bicomponent fibers can be split into two fibers as well.

**Segmented**

Various types of segmented fiber cross-sections are shown in Figure 7.2. Segmented-pie fibers are made of segments of two polymers as in a ‘pie-diagram’. The fibers with an 8-segmented pie or 16-segmented pie are the popular ones used as precursors for microfibers. Splitting apart the components of segmented-pie fibers produces microfibers that are finer than conventional microfibers, but these are mixtures of two polymeric microfibers. A hollow segmented-pie structure assists in splitting apart and generates finer microfibers. The cross-section that is most readily splittable into microfibers is a segmented ribbon. But they are difficult to card because of the anisotropic bending moment and very fine denier. To improve performance in the card, fibers with multiple lobes such as the cross and the trilobal cross-sections were tried. Generally, it is preferable to produce carded webs and then split the fibers.

**Islands-in-the-sea**

Islands-in-the-sea type bicomponent fibers are used to produce nanofibers using a novel method. Often the islands are made of PET, and the sea comprised of a special type of polyester that is hot water soluble or caustic soluble. Hydroentangling is one of the techniques used to break the islands apart from the sea. The separated islands form nanofibers, which are typically coarser than the electro-spun nanofibers.

**7.2.2 Non-circular cross-section and profile fibers**

Generic thermoplastic polymeric fibers have a solid circular cross-section. In the melt spinning process, the cross-sectional geometry of such fibers
Subjected to the heat transfer phenomenon that takes place in the quench chamber. The shrinkage forces act in the radial direction and lead to the circular cross-sectional geometry of the fiber. Other desired cross-sections such as hollow, trilobal, hexagonal, etc. (as shown in Figure 7.3) can be accomplished by using the desired spinneret-hole configuration. Non-circular capillary fabrication requires sophisticated electron beam milling and electro-discharge machinery. This involves additional capital cost for procurement. Furthermore cleaning, maintaining the profile and inspection of these spinnerets are more expensive. Hence non-circular cross-sectional fibers are produced when the value-added properties offset the additional cost involved.

Hollow fibers are lightweight and have excellent insulative properties. They are used as filler fibers in pillows and cushioned furniture. Trilobal cross-section fibers have reflective surfaces, giving rise to a sparkly appearance; hence they are often used in luxury apparel. Trilobal fibers provide rigidity and resilience, and the reflecting surfaces are efficient at scattering light to hide any dirt. For these reasons, trilobal fibers are often used in carpets. Similarly a ribbon cross section has flat surfaces for the reflection of light, so its appearance is also highly sparkly. Ribbon fibers pose much
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oped by Eastman Chemical Co., Kingsport, TN, and then donated to the Clemson University Research Foundation. The 4DG cross-section is shown in Figure 7.4 and other cross-sections with high shape factors are shown in Figure 7.5. The deep grooves or channels along the longitudinal axis of the fiber are unique features of the fiber that can help move fluid spontaneously, trap particles, provide large surface areas, and bulk cover per denier per filament. Usually, either hydrophilic or hydrophobic surface finish is applied to the fibers, based on the application. 4DG fibers can accumulate more dust particles if they are used in filters or if they carry more carbon particles placed in the grooves for odor absorbency on the right. 4DG is one among the various thermal insulation materials used in a Mars space suit. Presently Fiber Innovation Technology, Inc. Johnson City, TN manufactures the 4DG fiber. Coolmax fabrics produced by Invista consist of fibers with uniquely engineered micro-channel cross-sections with a denier gradient, but the channels are not deep as in 4DG. The fabrics move moisture away from the body, thus keeping the skin cool, dry, and comfortable. Potential applications are: perspiration shields, sweat bands for athletes, mascara brushes, condensate collectors, mops and brushes, socks, cosmetic applicators, oil absorbers, wipes, filter media, and wound care materials.

7.4 ‘4DG’ fiber cross-section with deep grooves.

7.5 Cross sections with high shape factor.
7.2.3 Monofilaments

Monofilaments are continuous filaments with high tenacity used in engineered fabrics or yarn. In order to achieve this, a high viscosity of the raw material is essential along with a higher level of purity. The conventional process produces PET polymer of intrinsic viscosity of about 0.65 dl/g maximum. An additional processing step, the solid-state polymerization (SSP) process, is required to upgrade the intrinsic viscosity from 0.6 to 1.0. Polycondensation is carried out by fluidizing the polyester chips in hot nitrogen gas at ∼210°C (about 10 to 20°C below the softening point of polyester). SSP is a very slow process, and it needs 20 to 30 hours of residence time in the reaction column, as the rate depends on removal of volatiles by diffusion through the polymer into the fluidizing gas (nitrogen). Depending on the application, newer polymers such as ultra-high molecular weight polyethylene (UHMWPE), polyethylene naphthalate (PEN), etc. are used to produce monofilaments with profiled cross-sections or chemically treated surfaces.

7.2.4 Microfibers and nanofibers

Fibers with a denier of less than one are known as micro-denier or microfibers. Generally, microfibers are produced by the melt blowing process. Such microfibers have diameters in the range of 1 to 15 microns. The melt blowing process is simple in machinery and construction, but a lot of variables are involved in determining the fiber diameter, fiber diameter distribution and the web structure. The process produces webs as individual fibers cannot be collected or handled, and these self bonded webs are used as filters for various applications. Almost any thermoplastic polymer can be melt blown to produce a microfiber nonwoven web.

Further, fine fibers with a diameter of less than 100 nm are called nanofibers, and usually are produced by an electrostatic spinning process, which is discussed in the following chapters. Recently, a lot of research has been done in the area of electrospinning as this process allows the production of nanofiber webs from several polymers, and the nanofiber structure has tremendous surface area. This makes these webs suitable for absorption, filtration and other related applications.

7.3 Production, properties and applications of chemically modified fibers

Specialty fibers can be produced from chemically modified polymers as well as copolymers. Modifiers are the chemicals added at some stage of the
polymer process or during extrusion. Table 7.1 contains the list of modifying chemicals, specifically chosen for the desired application.

### 7.3.1 Low pilling fibers

Pilling is the formation of little knots of fiber on the surface of a fabric, as a result of wear, contact, and abrasion. As a fabric rubs against another surface, fiber is pulled away from the yarn, protrudes, and entangles to form a knot on the surface of the fabric. This occurs in all fabrics and gives them a fuzzy appearance, more often in staple fiber based fabrics. On natural fabrics, when pills form they break and fall off. Since synthetic fibers are stronger, the pills formed do not break away, and hence they are more noticeable on synthetic fabrics than on natural fiber based fabrics. Chemical modification to achieve low pill polyester fibers involves the introduction of a weak spot in a strong polymeric chain. Generally, the modifier, glycerol or pentaerithritol, is added in the bulk polymerization. Other methods make use of microfibers or a low-pill finish, which looks excellent after wearing and laundering.

### 7.3.2 Improved dyeable fibers

Unlike natural fibers, polyester fibers are difficult to dye because of their chemical structure. Their dye pick up is very low, in spite of the use of high pressures and high temperatures. Modifiers like polyethylene glycols contribute slight openness in the polymer chain structure that eases dyeability. But in order to obtain darker and brighter shades comparable to silk, it is necessary to introduce dye reactive spots in the polymer chain.
The popularly used reactive modifiers contain compounds based on 5-sulfo-isophthalic acid; they are incorporated into the chain during the synthesis of polyester. As these modifiers are costly, the cationic dyeable polyester fibers and fabrics are expensive.

Nylon 6 has good acid dyeability. The dye pick up is due to the large number of amino acid (–NH$_2$) end groups in the polymer chain. Regular textile grade fibers have about 60 equivalent amino acid end groups. By controlling the quantity of amine stabilizers in the process, amino acid end groups could be brought up or down to alter the level of dye pick up. Generally, when amino acid end groups are above 100 equivalents, nylon is ultra-deep dyeable and when amino acid end groups drop below 15 equivalents, it dyes in light shades. Sometimes, in order to impart basic dyeability, anionic sulfonate (–SO$_3$) groups are introduced into the nylon chain.

7.3.3 Flame-retardant fibers

In order to obtain flame-retardant (FR) properties, about 3 to 5 wt. % phosphorus compounds are inserted into the polyester macromolecules. Kosa$^6$ produces an FR polyester fiber with the trade name Avora® using a proprietary organic phosphorus compound which is incorporated into the polyester polymer during polycondensation. Avora fabrics are used extensively in the hospitality industry, the healthcare industry, and public areas. Flame resistance is built into the fiber chemistry, such that the protection does not wash out, age out, and meets flammability standards. Due to the presence of the FR compound, the melting point is marginally lower than regular polyester. However, the compound makes it difficult for combustion to take place as the fiber shrinks away from flames, chars, reduces melting drips from occurring, and prevents further burning by self-extinguishing. Combustion products are similar to generic polyester comprising of oxides of carbon and water vapor, as well as small quantities of aldehydes and ketones. Recent innovations include compounding with aluminum oxides, nanoclays and nanosilicates to impart flame retardancy. The use of halides is banned due to environmental and health hazards. Polyamides, however, have proved difficult to render durable flame retardancy by incorporation of additives because of their melt reactivities. Other non-polyester FR fibers in the market include Basofil, Nomex, Kanecaron, and Visil.

7.3.4 Antistatic and conductive fibers

Antistatic consists of a conductive core or layer along the fiber length made of conductive carbon or metal. Nega-stat$^7$ bicomponent antistatic fibers
consist of a tribal shaped conductive carbon core and a polyester sheath. Resistat™ conductive fibers are produced by a carbon suffusion process that saturates the outer skin of the nylon fiber with electrically conductive carbon particles. The carbon practically becomes part of the structure of the fiber, which retains the strength and flexibility of nylon while maintaining excellent conductivity. A block copolymer containing polyether segments and a polyetherester also exhibits antistatic properties. Blending with polyether makes nylon fibers hydrophilic, and they absorb more moisture, which imparts an antistatic property. Antistatic fibers containing 5–15% conductive materials are blended with non-conductive fibers in stripe patterns to provide effective static dissipation for the life of the product. Antistatic fibers neutralize surface charges by induction and dissipate the charge by conduction when grounded or when ungrounded by air ionization, known as Corona discharge. Applications include industrial bags, brushes, conveyor belts, filters, carpets, and protective garments.

7.3.5 Biodegradable fibers

Polyester in its generic form, polyethylene terephthalate (PET), is not biodegradable. There are chemical modifications available to imbibe biodegradability. Biodegradable fibers are designed such that at the end of their useful life, they can be sent for composting, where they can decompose in an environmentally friendly way, which promotes the growth of plants, earthworms and microbes in the composting soil. Eastar® is a thermoplastic and biodegradable binder fiber produced by Eastman Chemical Company. Eastar is co-polyester with melting point ∼120°C. Eastar is easily bondable with cellulosics, and the finished product is completely biodegradable. Biomax® from DuPont is another biodegradable polyester fiber. Biomax is modified hydro-biodegradable polyester with a melting point of 200°C. Proprietary monomers are incorporated into this polymer, creating sites that are susceptible to cleavage by moisture into smaller molecules, which are then consumed by naturally occurring microorganisms and converted to carbon dioxide, water and biomass.

Polyactic acid (PLA)® from Dow-Cargill is a biodegradable polymer that is produced from cornstarch. PLA fiber has a melting temperature of 175°C and tensile properties comparable to that of polyester fibers. PLA, as a melt spinnable fiber from a vegetable source, has many of the advantages of both synthetic and natural fibers. In addition to being produced from a renewable raw material, it is biodegradable. However, PLA has poor abrasion resistance and bonding behavior when compared to conventional binders. Other biodegradable fibers are Ecoflex® from BASF and polyhydroxybutyrate-valerate (PHBV) from Metabolix.®
7.3.6 Low melt binder fibers

The melting point of the most commonly used polyester fiber, polyethylene terephthalate, is about 260°C, but it can be reduced through the incorporation of modifiers such as isophthalates, adipates, etc. Polyester with a melting point of ~245°C is a commodity polymer used in making polyester bottles for soft drinks. These modifiers are added during polymerization, and they cause some distortion and defects in the polymer chain leading to a drop in the melting point accompanied by a minor drop in the glass transition temperature. A drop of about 2 to 3°C in the melting point is obtained by a percentage addition of the isophthalate group in the polymer chain. Adipates cause very large drops in the melting point, two to three times that of isophthalates. Low melt fibers (with melting point of 110°C or 130°C) are usually blended with commodity fibers in the desired proportion and then thermal bonded into webs. Generally bicomponent fibers with a low melt polyester sheath and a conventional polyester core are popular in the market. The binder fiber fuses with other non-melting fibers in the vicinity at the contact point, and it preserves the integrity of the nonwoven. A major advantage is that it creates a fabric free of chemical bonding agents.

7.3.7 Caustic soluble fibers

The presence of the right proportion of long chain polyethylene glycols, sodium salts of sulfo-isophthalate compounds, etc. helps to imbibe caustic solubility in polyesters. A novel method of producing microfibers involves the dissolution of the sea in the ‘islands-in-sea bicomponent fibers’ wherein the islands are made of conventional PET, and the sea is made of caustic-soluble polyester.

7.3.8 Chemical and heat resistant fibers

Increased chemical and heat resistance is achieved by the introduction of bulky groups such as naphthalenes and butylenes in the polymer chain. One-hundred percent polyethylene naphthalate (PEN) has high glass transition temperature (125°C) compared to PET (68°C). Moreover, PEN fibers start to shrink significantly at 190°C [150°C for PET]. Naphthalene molecules in the polymer chain impart higher tensile strength, UV stability, and high temperature resistance. They also have better barrier properties and are more resistant to hydrolysis in alkaline or very hot aqueous conditions. PEN fibers are used in sail cloths, and hot liquid filtration media. However, they are more expensive and have a lower flex life. To lower the
cost PEN blends\textsuperscript{14} with PET have been tried. Polyphenylene sulfide (PPS)\textsuperscript{15} and Halar\textsuperscript{16} fibers are also used in such applications.

### 7.3.9 Water absorptive fiber

Water absorptive fibers\textsuperscript{17} are used in sportswear to absorb and take away the sweat. Hollow polyester fibers have porous walls with pore size 10 to 30 nm in diameter and absorb water by a capillary phenomenon. The pores run through the surface into the hollow core. QUUP is water absorbing nylon fiber that contains proprietary nitrogen based polymer that has water absorb/release properties. HYGRA is another nylon skin-core fiber where core is made up of hydrophilic polymer.

### 7.3.10 Deodorant fiber

Deodorant fiber has the ability to absorb volatile compounds from the atmosphere, generally tested with formaldehyde. Daiwabo has developed polymer with proprietary amine compounds in the chain that absorbs formaldehyde as well as other vapors such as ammonia, acetic acid, and cigarette smoke.

### 7.3.11 Smart fibers

Smart\textsuperscript{18} fibers for different applications are made from chosen polymers for specific applications, and they are different from generic polyesters or polyamides. Smart fibers are generally tailored for specific applications such as artificial muscles, intelligent biomedical garments, adaptive textile structures, thermo-regulated snow clothing, etc.

### 7.3.12 Micro grooved fibers

Micro grooves on fibers are obtained by blending micron sized additives while producing the fibers and then taking away the additives at the finishing stage, mostly by dissolution. While drawing, stretching, and setting the blended additives do stretch and leave groves along the fiber at the final stage.

### 7.3.13 Lotus leaf effect

Water and soil repellency in fibers is achieved by imparting ultra hydrophobic nature through lotus leaf effect.\textsuperscript{19} Basically the fibers need to have low surface energy and have the extended degree of surface roughness. A combination of a surface attached polymer layer and nanoparticles provide
the desired results. A polystyrene layer is grafted onto polyester fiber. Further the topography and the resulting roughness of the surface is tuned, by changing the size and shape of the nanoparticles. Ultimately the water contact angle close to 130° is found to provide adequate hydrophobic properties.

7.3.14 Surface treated fiber – dull luster

In order to control luster, generally titanium dioxide is added at the polycondensation stage of the manufacture. Fiber surface treatment by chemical means is a method to modify the characteristics of smooth fibers as it imparts dull luster as well as new textile properties such as fluffiness and warmth retention. Generally caustic chemicals are employed in the treatment.

7.3.15 Fibers with controlled shrinkage

Heat setting is the operation carried out to control the residual shrinkage in the fiber. Generally fiber is subjected to heat under tension by passing over oil heated drums. In this process as the crystallinity increases the shrinkage reduces. Fibers with low shrinkage (less than 1%) are used in sewing threads. Other approaches such as adding nucleating agents to enhance crystallization during fiber formation are studied extensively but their commercialization is limited.

7.3.16 Other specialty fibers

Compounding with nanoclays or nanosilicates enhances mechanical properties such as heat distortion temperature and tensile strength as well as the flame retardancy of the product. Nanoparticles have a very high aspect ratio and hence possess a high surface to volume ratio. In order to obtain the advantages, it is essential to have proper surface wetting, dispersion and exfoliation of the nanoparticles. The polymer-layered silicate (PLS) exhibited increased modulus and strength, decreased gas permeability, and decreased flammability. While nanoclay adds muscle to plastics, carbon nanotubes impart electrical and thermal conductivity. Single wall and multiwall carbon nanotubes are relatively new, and their commercial potential has been limited by their high prices.

7.4 Design and process control aspects

Modifiers and additives cause some changes in specialty polymer behaviors with respect to heat transfer, glass transition, melting, crystallization, and
orientation of the molecular chains compared to the conventional polymers, and hence call for some minor adjustments in the processing conditions. Basics of fiber technology\textsuperscript{23} and advanced fiber spinning\textsuperscript{24} involve process as well as heat transfer parameters. Conventional polyester fibers are made from polymers having an intrinsic viscosity (IV) of 0.600 to 0.650. For specialty fibers, more or less the same melt viscosity is required, but chemical modification can offset the relation between melt viscosity and IV. In certain specialty fibers, the process-loss in viscosity during melting and spinning is more than normal (~0.020). In such cases, an appropriate adjustment is to go for slightly higher viscosity polymer granules. Specialty fibers with modified fiber cross-sections are obtained by the controlled solidification of the melt streams emerging out of the spinneret capillaries. The final cross-section depends upon the machine and process variables.

Operating variables are online variables that can be changed while the machine is being operated. These variables include throughput and temperatures of the polymer and quench air, collector or winder speed. The quench air is adjusted towards a lower velocity profile for finer fibers. There are a few variables at the drawing and heat setting stages. All of these affect the final properties of the fiber. Off-line variables are fixed before starting the spinning process, and can only be changed with downtime or when the machine is not in operation. These variables include spinnerets (capillary design) and quench air profile setting hardware.

In the case of conventional fibers, the spinneret holes or capillaries are in the diameter range of 0.1 to 0.5 mm, with a length to diameter (L/D) ratio of 2 to 5. Finer filaments are produced with the smaller diameter capillaries, at lower melt speeds and at slightly higher melt viscosity. Melt extrusion velocity is in the range of 5 to 75 m/min (for conventional fibers 5 to 50 m/min).

For specialty cross-sections, calculations are carried out based on the same principles used in conventional fibers with a slightly more complex formula derived for the specific geometry. In the case of hollow fibers, in order to get a fiber cross-sectional inner diameter to outer diameter ratio (d/D) of 0.3, spinneret capillaries d/D of about 0.6 are required. Generally, the capillary bridge width is about 20% of the outer diameter of the capillary. In the case of ribbon fibers, in order to get a fiber cross-sectional length to width ratio (L/w) of 4, it is required to have spinneret capillaries L/w of about 12. For trilobal fibers, the preferred cross-sectional leg length to width ratio of the spinneret capillaries (L/w) is 3.

### 7.5 Future trends

With the continuing pressure for better quality, assured performance at lower prices is the primary goal of a manufacturer of commodity products.
Research interests often go for niche products in relation to the specific performance properties. This encourages the search for new materials and innovative approaches to satisfy the short-term and long-term goals. Composites containing polymers, fibers, nonwovens, films, and foams are growing and demonstrating their potential to replace conventional materials. The coming years will see dramatic changes in the level of their acceptance and the diversity of their use, as engineers gain control over the necessary design skills, and wider adoption makes them more cost effective. Moreover, with increased thrust for using raw materials from renewable resources, biodegradability of the product is demanding a shift from conventional synthetic fibers that are too stable and do not degrade in the municipal waste.

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8

Property enhancement through blending

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8.1 Introduction

The increase in the production of synthetic fibres in the past few decades has been phenomenal. This can be broadly attributed to two factors. First, lack of availability of natural fibres and second, controllable and superior properties of synthetic fibres like polyester and polyamide. These synthetic fibres have many superior properties such as uniformity in length and strength, freedom from foreign impurities and the flexibility in regard to the desired staple length and denier. In addition, some highly popular fibres like polyester impart higher tenacity, greater abrasion resistance and better crease resistance to the fabrics. However, these fibres have some negative attributes like proneness to static generation, low moisture regain, etc. Therefore, for most of the end uses especially the apparels, the common practice in the textile industry has been to use blended yarns instead of 100% single fibre yarns. Blending of fibres is also carried out in technical uses as well to enhance the properties of the final product or to reduce the cost of the product.

The aspect of blending of polyester and polyamide fibres with other fibres can be considered in two different ways: one, for apparel applications and two for production of speciality products like core spun yarns and nonwovens and other related products. For apparel end uses, blending is normally carried out in the fibre stage, i.e, blending of different fibres in the staple form and spinning them into yarn. In the case of core spun yarns, blending of one component in continuous filament form and the other in the staple fibre form are combined at the yarn spinning stage. In nonwoven processes, blending of different types of staple fibres or already formed yarns is carried out in one of the nonwoven processes like needle punching or stitch bonding to obtain a blended product. The majority of the polyester blends go into making the apparels where yarn uniformity in terms of mass variation and dyeability are critical. Hence, in this chapter the principles and methods involved in blending of staple fibres, particularly with
polyester and nylon, are discussed first. Second, significant amounts of polyester and polyamides are combined with other fibres to make industrial clothing, conveyor belting, tentage, elastic yarns and sewing threads through core-spinning technology. Hence, in the second section of the chapter, the technology of core spinning and other technical textile products made through nonwoven processes are discussed.

**8.2 Staple fibre blending**

The blends normally consist of two (binary blend) or three different fibres (tertiary blend). Such blending of two or more dissimilar fibres offers an effective means of projecting the positive attributes or aspect of each of the constituent fibres. Blending, however, needs to be carried out in such a way that the fibres in a blended yarn are thoroughly dispersed and intermingled. Any deficiency in proper blending of fibres results not only in poor yarn properties but also fabric defects like fabric streakiness when dyed.

All fibres, whether natural or manmade, are quite in-homogeneous in respect of their characteristics. Blending, therefore, is a must to bring the desired homogeneity in their characteristics even in a 100% single fibre stock. Blending of dissimilar fibres is undertaken to achieve some other important objectives too and is quite common in the industry.¹

In the case of polyester and polyamide fibres, blending is performed mainly in order to²:

- Give the required characteristics to the product (e.g. blending of polyester fibres with natural fibres produces the desire easy-care character).
- Hold on raw material cost (blending in of relatively cheap raw material).
- Influence favourably the behaviour of the material during processing (use of segmented fibres having alternate segments of polyester and nylon).

The major objectives of blending are improvement in functional properties, improved process performance, better economy and improved aesthetics. One of the primary reasons for blending is to improve the functional properties of the fabrics such as higher tensile strength, uniformity, better appearance, increased wear life, crease resistance, crease recovery, dimensional stability, elasticity and comfort. A 100% single fibre yarn cannot possibly impart all of these desirable properties to the fabric. Take the case of polyester–viscose, a very popular blend. Fabrics from 100% viscose rayon suffer from low tensile and tear strength, poor crease resistance and recovery and low abrasion resistance. However, addition of polyester can
overcome these deficiencies because it has an excellent dimensional stability, crease retention, crease recovery, quick drying, increased abrasion resistance, higher tensile and tear strength. However, 100% polyester fabrics are also not desirable as they are prone to static accumulation, hole melting and pilling. They are moisture resistant (hydrophobic), difficult and expensive to dye and have a poor hand. Like polyester and viscose, each fibre has its own positive and negative attributes. These negative attributes of polyester and viscose can be reasonably neutralized by adding a certain percentage of each fibre. For example, the addition of viscose imparts to the polyester fabric a reduced static propensity and a marked protection against pilling. This also results in increased fabric absorbency and wear comfort. The other gains accruing from such a blend are the increased washability, increased affinity for dyestuffs and chemicals used in finishing.

Some fibres like polyester at times are quite troublesome to process in 100% form especially at the card. Addition of fibres like cotton or viscose rayon in the previous process has been seen to facilitate the smooth carding of such fibres. The blending of synthetic fibres which are longer and finer than cotton influences the spinnability as well as the productivity. Addition of a certain proportion of long fibres favourably influences the drafting characteristics of short staple fibres by acting as carrier fibres. The addition of fine fibres to a blend increases the number of fibres in the cross-section of a given count of yarn which leads to extension of the spinning limit. The longer fibres in the blend, in the same way, make it possible to spin finer counts. In ring spinning, such blending of longer and finer fibres gives substantial increase in production through a reduction in the required twist level, lower end breakage rate and higher spindle speeds. Such advantages obtained at the spinning stage seem to extend to a large extent to every post-spinning operation up to weaving or knitting.

Fabrics with a variety of colour mixture or shades can be produced by blending differently dyed fibres at the blowroom, drawframe or roving stage. Use of various fibres with different length, denier, colour and surface characteristics are employed to obtain unique fabric surface by exploiting the varying drafting and migratory behaviours of the fibres. Such effects are used to enhance the aesthetic appeal of a fabric.

Aesthetics of the fabric and garment are acquiring greater influence on the purchase pattern. The important aspects are fashion, texture, drape and lustre. All these attributes can be manipulated through selection of right type of fibre characteristics and specifications. The aesthetics of a fabric can also be developed by selecting specific blend components and their proportions. These parameters can be exploited to impart increased bulk, an attractive raised finish or a pleasing total variation to the fabric.

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8.3 Evaluation of the blend

The evenness of the blend must always be assessed in two directions: the longitudinal direction and the transverse direction. Where there is unevenness in the longitudinal direction, successive yarn portions exhibit different percentage distributions of the individual components (Figure 8.1). These can lead to stripiness.

Where there is unevenness in the transverse direction, the fibres are poorly distributed in the yarn section (Figure 8.2). This irregularity leads to an uneven appearance of the finished product.

The determination of the evenness of a blend, e.g. of synthetic and natural fibres, is costly and not simple. One component is usually dissolved out or coloured differently.

8.3.1 Indices of blending

For optimum exploitation and reasonable prediction of yarn and fabric properties from the properties of the constituent fibres, it is essential to
achieve a homogeneous or an intimate blend. Ideally, the blend should be a perfect blend in which every fibre property like length, denier and tenacity, etc. is found in the same proportion within every unit cross-section of the yarn. However, such a blend can never be achieved in practice. The best one can achieve, is the one in which the constituent fibres of the blend are randomly distributed throughout the cross-section along the length of the yarn. This random distribution is seen to be close to a Poisson distribution. Generally, it is easier to obtain satisfactory results in this regard in a 50/50 blend than in an 80/20 blend. One of the important factors in achieving an intimate blend is to achieve the smallest fibre aggregates to the extent possible at the earliest stage of spinning. The blend homogeneity can be expressed either as the Degree of Mixing or the Index of Blend Irregularity.

**Degree of mixing**

This is a statistical parameter, which estimates the inherent intimacy of a blend. DeBarr and Walker\(^3\) made a series of yarns from a blend of black and white fibres and examined their cross-sections for the number of groups of fibres. They assumed that the fibre distribution in yarn sections would become random as the number of doublings approached \(\infty\). If \(g^∞\) is the number of groups of white fibres, one can write:

\[
g^∞ = np (1 - p)
\]

where \(n\) is the number of fibres in the average cross-section of the yarn and \(p\) is the average proportion of white fibres in the blend.

In practice, the number of groups of white fibres is less than this since the number of doublings in use never achieves perfect randomization. If \(g\) represents the actual number of groups of white fibres, the degree of mixing \((\gamma)\) is given as,

\[
\gamma = \frac{g}{g^∞}
\]

A value of 1 for \(\gamma\) shows a thorough random distribution of fibres. A value of less than 1 means less than random mixing and a value of more than 1 means that the mixing is better than random. As this value increases, the blend approaches a perfect blend. In practice, the value of \(\gamma\) is less than 1 as the number of doublings increases. The finer yarns, therefore, require fewer doublings to achieve random distribution.

**Index of blend irregularity (IBI)**

This is another statistical measure used to assess the degree of randomness of fibre distribution. This index was developed by Coplan and Klein\(^4\) for...
variation in the blend proportion against the theoretical value for random mixing. The IBI can be calculated from the following expression,

$$IBI = \sqrt{\frac{1}{M} \sum_{i} \frac{(T_i p - W_i)^2}{T_i p q}}$$

where

$T_i$ is the total number of fibres in a given cross-section,

$W_i$ is the number of fibres of component $W$ at the cross section,

$p$ is the average fraction of component $W$ for all cross-sections,

$q$ is equal to $(1 - p)$ and

$M$ is the number of cross-sections examined.

Obviously, this index is primarily a chi-square test. It is a normalized standard deviation. A value of zero would mean perfect blending and would indicate random blending. Values greater than 1 indicate less homogeneity than what complete randomness would give. This index can also be used to calculate the fibre cluster size. An index of 1.3 has the physical meaning that the blend suffers from causative factors that increase the homogeneity by 30% over what can be expected from a purely random process operating on single-fibre elements. By considering fibres to be in clusters, replacing $T_i$ by $T_i/C$, where $C$ is cluster size (number of fibres in the cluster), and substituting 1 for $IBI$, an estimate of cluster size can be obtained. Experiments on wool/nylon yarns spun on the woollen system suggested a cluster size of 2–4 fibres. Coplan and Bloch expanded on the longitudinal variation method and developed two other methods of evaluation namely, radial distribution and rotational distribution of fibres.

Radial distribution

This describes the fibre motion across the cross-sections of the yarn. To estimate this, the yarn cross-section is divided into, usually four, concentric circles of either equal area or thickness. Figure 8.3 shows circles of equal
thickness along with the blend ratio plotted in the form of a bar diagram. Ideally, the blend ratio should be the same in all four zones from inside to outside.

Rotational distribution

For estimating the variation in the rotational distribution, the yarn cross-section is divided into four to six segments. The blend ratio is calculated and plotted as bar diagram as shown in Figure 8.4. Ideally the ratio should be the same for all the segments.

8.4 Migration

The segregation of components in a blend yarn is known as migration. A comprehensive understanding of this aspect of fibre behaviour is essential for the proper selection of fibres for a blend. The migratory behaviour of fibres in yarns arises from differences in tensions developed during twisting of fibres into yarn. The fibres twisting around a long path on the yarn surface would develop a high tension while fibres following a shorter straight path would be under lower tension. The tension differences would cause an interchange of position between the high and low tension fibres and thus lead to migration. It is known that, in a blend yarn the fibres having higher modulus tend to occupy the inner zones of the yarn cross-section and those fibres having lower modulus, the outer zone.

Cassie postulated that for any spun yarn, the tendency for fibres to coil up and pack closely when twisted, depends on the parameter, \( \frac{1}{h} - \frac{2}{E} \), where \( h \) is the modulus of torsional rigidity of a fibre and \( E \) is its bending modulus. The greater the positive value of this parameter, the closer is the packing. Therefore, in a binary blend the component with a greater value of \( \frac{1}{h} - \frac{2}{E} \) will pack more tightly and hence tend to occupy the core of the yarns. Obviously, the component with the lower value of \( \frac{1}{h} - \frac{2}{E} \)
will be displaced to the surface. In general, longer, finer and higher modulus fibres tend to migrate to the core of the yarn.

The dyed fibres and non-crimped fibres also tend to migrate to the core of the yarn. Therefore, in a blend yarn having dyed and undyed fibres or crimped and uncrimped fibres, the undyed and crimped fibres will predominate on the surface of the yarn. Fibre migration has significant influence on the performance and functional properties of yarns and fabrics. The migration is essential for producing a strong yarn by providing interlocking between different fibre layers of the yarn cross-section. A yarn with ideal helical geometry will not have such interlocking between these radial layers and there will be nothing to stop the fibres on the surface peeling off so that the whole yarn can be easily rubbed away.

In a binary blend if the stronger component migrates preferably to the core, the resultant blend yarn will have a higher tenacity. The fabric properties which are affected by migration include the pilling tendency, the soiling, the handle or feel, the drape and the abrasion resistance of the fabric. The defects like soiling and pilling can be contained by judiciously choosing the fibre properties of the blend components so as to put them at desired locations in the yarn cross-section. The extent to which this objective is achieved can be evaluated by determining the migration index.¹

### 8.4.1 Migration index

The migration index of a blend can be calculated by using Hamilton’s method.⁷ This index is based on the calculation of certain first moments of the blend component about the centre of the yarn cross-section and relates the moment $FM_a$ corresponding to the actual distribution, to the moments $FM_u$, $FM_i$ and $FM_o$ corresponding to three hypothetical distributions viz., uniform distribution, and those which would result from maximum inward and outward migrations, respectively. This is illustrated in Figure 8.5.

$FM_o - FM_u$ represents maximum possible outward migration.

For a polyester-cotton blend the expressions used for the calculation of this index are,

\[
\text{Migration index for polyester} = \left( \frac{(FM_a - FM_u)}{(FM_u - FM_i)} \right) \times 100
\]

\[
\text{Migration index for cotton} = \left( \frac{(FM_a - FM_u)}{(FM_o - FM_u)} \right) \times 100
\]

The above expressions are based on the fact that the polyester is going to migrate to the core and the cotton towards the surface. The two indices

\[
FM_i \quad \quad FM_u \quad FM_a \quad FM_o
\]

8.5 $FM_a - FM_u$ represents actual outward migration.
will be seen to be equal but have opposite signs, i.e., if one is positive, the other will be negative. A negative value for a component indicates that it has preferentially inward migration whereas a positive value indicates outward migration.

8.5 De-blending

The spinner aims to distribute the different fibres evenly in the yarn. For this purpose he must first produce a good blend at some stage of the process, and second be able to maintain the blend up to the stage of binding into the yarn. As is well known, meeting the first of these requirements is not always easy; sometimes the second is harder still. Fibres of different length, different surface structure, crimp, etc., also behave differently during movements as individuals. A ‘de-blending’ effect can very often arise.

During rolling of fibres in hoppers (bale openers, hopper feeders), migration of the components occurs where the fibres have different structures (e.g. cotton and synthetic fibres). A similar effect is found during drawing in drafting arrangements. Fibres with large differences in length or surface form (smooth/rough, dyed/undyed, etc.) do not exhibit the same cohesive contact with each other. When a drafting force is applied, they move differently – this leads to clumping of fibres into clusters and finally to de-blending. Pneumatic transport can also cause de-blending.

8.6 Selection of blend constituents

The choice of blend constituents needs careful consideration both in terms of the fibre type and their proportion in the blend so as to bring out their positive attributes and to suppress the negative ones. One has to further decide the specifications of these fibres, keeping in mind the end use requirements. For example, one may have to choose between low tenacity and high tenacity polyester, round cross-section or trilobal polyester.

8.6.1 Type of fibre

The major objective of blending two or more fibres is to produce a yarn or fabric with improved functional properties, which a single fibre just cannot achieve. Take the case of some popular blends like polyester-cotton or polyester-viscose, which are normally used, for apparels like shirting and suiting. If the fabric is made from 100% polyester it will have excellent crease resistance, durability, and wash and wear due to the hydrophobic nature of the polyester fibre.
Addition of a hydrophilic fibre like cotton or viscose rayon may render it more comfortable to wear because of their higher thermal conductivity and better absorption of perspiration. The body heat and perspiration are thus easily passed on to the outside air resulting in wear comfort. Such a blend fabric, therefore, overcomes the clammy feel of the 100% synthetic-fibre fabric. The fabric would also save the user from the pricking sensation due to lower bending rigidity of the cellulosic fibres and thus give it a softer feel.

The choice between cotton and viscose rayon for blending with polyester is primarily dependent on the end product. For a fuller look in lightweight fabrics, such as shirting, cotton is preferred. Even for heavier sorts such as working clothes and dyed uniform fabrics, cotton may be preferred to viscose so as to avoid a differential shade at the worn out portions. The viscose rayon fibre due to its lower abrasion wears out much faster than cotton, thus giving prominence to shade of polyester fibres which is generally somewhat different from that of viscose fibres. The blend with viscose would, however, give superior results to cotton in terms of yarn evenness, appearance, freedom from trash particles and contaminants. This blend is, therefore, preferred for medium and heavy constructions, such as suiting, where impeccable appearance is important. Similarly, blending of shrinkable and non-shrinkable polyester or other type of fibres is carried out to produce hi-bulk yarns.

The blends of two fibres, called binary blends, account for the major portion of the blended yarns produced. However, to a limited extent, the tertiary blends (blend of three types of fibres) are also produced. One such blend is the polyester-cotton-polynosic. Such blends are normally used for reasons of economy, styling and special surface effect in the fabrics. The objective of exploiting the positive attributes of each of the fibres is relegated to a secondary place in such tertiary blends. More often than not, such blends are produced for commercial exploitation by carrying the name of the most desirable but costly fibre though it may be present only in a small proportion. The tertiary blends can also be used for production of fabrics having multi-coloured effects during piece dyeing. This, however, demands a great skill on the part of the dyer and a thorough knowledge of the reactivity of one group of fibres with another.

8.6.2 Compatibility of fibre blends

For trouble-free processing and optimum utilization of the positive attributes of the blend constituents, it is imperative to consider their compatibility in terms of their properties such as length, denier, breaking elongation, etc.
Length and denier

The two most important fibre properties, which need to be matched for satisfactory processing of blends, are the staple length and the fineness of the fibres. As a general rule, these two fibre properties should be nearly the same for all the constituents unless deviations are desired for producing some special effects. For blending of polyester with combed cotton of, say 34 mm, the denier of cotton component, generally in the range of 1.3 to 1.5 (3.6 to 4.2 μg/inch), is matched with 1.2 to 1.5 denier of polyester. Similarly in a viscose-cotton blend the viscose rayon of 1.5 denier and 29–32 mm length is generally used to match cotton of 1.5 denier (4.2 μg/inch) and 28 mm length.

The denier difference, on the other hand, seems to be detrimental to yarn quality as the migratory behaviour during twisting in yarn formation prevents intimate blending of fibre during twisting; the coarser fibres going to the surface and the finer to core. Excessive difference in the denier of the blend fibres, say 1.5 denier viscose and 3.0 denier polyester, has been seen to produce very weak yarns. It is difficult to lay down any firm tolerance limits for the difference in length and denier of the blend fibres. In general, it can be stated that if the differences in the length and fineness are high, the process performance and the yarn quality are expected to be poor.

Extensibility

The elongation compatibility of the constituent fibres is the most important fibre property in respect of achieving the maximum strength from a blend. In a blend, the fibre with a lower breaking elongation will break when the other fibre component is sharing only a part of its breaking load. After the breaking of the less extensible fibres, the full load is transferred to the more extensible component, as the less extensible fibres can no longer share any load. This extent of inequality in sharing of tensile load is determined by the load elongation curves of the two fibres.

Dyeing properties

In case the blend yarn or fabric is to be dyed subsequently, due consideration should also be given to the dyeing properties of the individual fibre components.

Tensile specifications of fibres

Having decided the type of blend fibres and their physical specifications, the next step involves careful consideration of the specific tensile
Polyesters and polyamides

choice may have to be made between low-, medium-, and high-tenacity tailor-made for blending with cotton. It be noted that the stress–strain curve for this type of polyester is quite different from the normal polyester. This is high tenacity polyester, sometimes called cotton type polyester. An exercise by Ranganathan and Vengsarker\(^8\) with viscose rayon as the second component has shown that the stress–strain curves of viscose rayon, given in Figure 8.7, closely follow that of medium-tenacity polyester.

The compatibility achieved in the aforementioned way optimized the yarn tenacity only. There would be many other situations where the matching is for a different end result. If the end product for a polyester-cotton blend happens to be shirting cloth, the hand of the fabric may take precedence over tenacity. It has been shown that these parameters can be considered in a quantitative way as these are well correlated with the compliance ratio. This ratio can be easily calculated from the stress–strain diagram of a fibre by using the following expression:

\[
\text{Compliance ratio} = \frac{\text{Maximum stress}}{\text{Maximum strain}}
\]

Figure 8.6 depicts such a matching of cotton and polyester fibres. It may be noted that the stress–strain curve for this type of polyester is quite different from the normal polyester. This is high tenacity polyester, sometimes called cotton type polyester. An exercise by Ranganathan and Vengsarker\(^8\) with viscose rayon as the second component has shown that the stress–strain curves of viscose rayon, given in Figure 8.7, closely follow that of medium-tenacity polyester.

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\[
\text{Compliance ratio} = \frac{\text{Maximum stress}}{\text{Maximum strain}}
\]

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Compliance ratio = \frac{10S_{10}^{-1} - 5S_{5}^{-1}}{5}

where \( S_{10} \) and \( S_{5} \) are the stress values in g/den at 10% and 5% fibre extension. Using this criterion the medium tenacity polyester is at the top of the category of moderate compliance ratio and high tenacity is at the bottom. Therefore, from the point of view of hand of the fabric, the obvious choice would be the medium tenacity polyester. The fact that the dye uptake of high tenacity polyester is less than that of medium tenacity fibre, the medium tenacity fibre may be preferred for general apparels. However, worldwide the blending of cotton is generally done with high tenacity polyester (6.2–6.4 g/den).

8.7 Blend ratio

The blend ratio is generally calculated on the basis of conditioned weight (20 ± 2°C and 65 ± 2% RH in tropical countries, 27°C may be the secondary standard). Having decided the type of fibres and their specifications, the next critical decision to be made is in regard to the proportion of each type of fibre, i.e. the blend ratio, so as to optimize the desired properties in the fabric. Extensive research has been carried out on this aspect with various types of fibre blends.

Figure 8.8 gives a generalized view of results obtained in these studies. These are given in graphical form for various fabric properties such as tensile strength, tear strength, crease recovery, crease resistance, abrasion.
2.32 Polyesters and polyamides

<table>
<thead>
<tr>
<th>Blend Ratio</th>
<th>Polyester</th>
<th>Polyester viscose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyester</td>
<td>67/33</td>
<td>67/33</td>
</tr>
<tr>
<td>Second fibre</td>
<td>0 - 25 - 50 - 75 - 100</td>
<td>0 - 25 - 50 - 75 - 100</td>
</tr>
</tbody>
</table>

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resistance, pilling, anti-static resistance, etc. for polyester-cotton and polyester-viscose fabrics. This figure can be used as a guideline for achieving the optimized fabric properties by selecting the proper blend ratio. It needs to be mentioned here that the yarn and the fabric construction as well as the final finishing of the fabric, especially with regard to heat setting and resin finishing, does make a noticeable difference in the desirable blend ratio. Another factor that influences the blend ratio is the stress–strain behaviour of the component fibres. In a polyester-viscose blend if the stress–strain curves of the two fibres do not match, the blend yarn is weaker at a low ratio of polyester. The optimum blend ratio would then work out to 65:35 for optimum wear performance. The polyester compo-
nent in the blend can be reduced to 50% if the fibres are made compatible in this respect. Some economic considerations, more than anything else, dictate the decisions about blend ratios that might otherwise look unreasonable.

### 8.8 Types of blending operation

#### 8.8.1 Possibilities

Blending of staple fibres can be carried out at various process stages, using various methods, equipment, machines and intermediate products. The following can be distinguished:

<table>
<thead>
<tr>
<th>Blending type</th>
<th>Process stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bale mixing</td>
<td>Before the blowroom</td>
</tr>
<tr>
<td>Flock blending</td>
<td>Within the blowroom</td>
</tr>
<tr>
<td>Lap blending</td>
<td>Using scutchers</td>
</tr>
<tr>
<td>Web blending</td>
<td>At the ribbon lap machine or the blending drawframe</td>
</tr>
<tr>
<td>Sliver blending</td>
<td>At the drawframe, the sliver-lap machine, or the comber</td>
</tr>
<tr>
<td>Fibre blending</td>
<td>At the card or the OE spinning</td>
</tr>
<tr>
<td>Roving blending</td>
<td>At the ring spinning machine</td>
</tr>
</tbody>
</table>

In addition, controlled and uncontrolled blending must also be distinguished. In uncontrolled blending, the components are brought together at random and without a mixing system (e.g. often in bale mixing). In controlled blending, the individual components are supplied to the machines in an ordered fashion and precisely metered (e.g. in weighing hopper feeders).

The various blending processes often differ strongly from one another with respect to capital cost, labour intensiveness, precision of blending, liability to error and simplicity. Each method has advantages and disadvantages. It is therefore not possible to put forward patent recipes for the use of one or another blending principle. The actual techniques used for blending of fibres can be referred to in the standard text books.

#### 8.8.2 Optimum blending method

The objective of choosing a specific blending stage is to achieve the most homogeneous blend consistent with process performance, yarn quality and fibre waste. The most important factor influencing the choice of blending method is the type of fibres to be blended. In case the component fibres need similar opening and cleaning treatment, the blending may be resorted
to at the blowroom, and that, too, at the initial stages through stack or machine blending. This stage of blending is most suitable for blends like polyester-viscose as both these fibres are free from foreign impurities and trash and require mild opening action. The stack blending or sandwich method is simple and is generally used for blending of man-made fibres among themselves. The rayon component, which is rather more cohesive, may have to be pre-opened before blending.

Polyester and cotton, therefore, is normally blended at the drawframe. In certain instances, however, the mills using drawframe blending for polyester-cotton face serious problems in the carding of 100% polyester, and dye shade variations in the blend fabric due to poor homogeneity of the blend. Such mills can advantageously use blowroom blending so as to overcome these problems. The presence of cotton fibre along with polyester would avoid any difficulty faced in carding of 100% polyester fibre because the cotton fibres act as carrier fibres for the polyester fibres. The improved blend homogeneity obviates the problem of dye shade variation to a large extent. Blowroom blending would, however, require opened and cleaned cotton. Therefore, the polyester stock is blended with combed cotton sliver through stack blending. This practice obviously involves processing of cotton component twice through the blowroom and the card. Such overprocessing of cotton might result in more neps in the blend yarn. Blending drawframe would obviate the need for this practice, if poor blend homogeneity were to be the only problem. As mentioned earlier, the blending drawframe gives the same level of blend homogeneity as the blowroom blending.

8.9 Influence of fibre properties and blend ratio on yarn properties

8.9.1 Tenacity

The tenacity of blend yarns has been studied by many research workers. Invariably, it has been found that the yarn strength of blend yarn is lower than the weighted strength calculated from the respective strengths and proportions of the component fibres. One of the more logical and simple procedures was put forward by Hamburger\textsuperscript{1,9} to predict the yarn tenacity with reasonable accuracy. This analysis has been represented graphically in Figure 8.9 for two components A and B for various possible stress–strain diagrams. The left-hand side shows the stress–strain behaviour of fibres and the right hand side the yarn tenacity for various blend proportions.

In all the three cases depicted in Figure 8.9, component A has lower breaking extension than component B. When a blended yarn is extended by applying a load, the fibres of the less extensible component A are
8.9 Effect of stress–strain diagrams of fibres on yarn tenacity with varying blend ratio.

strained to the breaking point first. This can be considered as the first rupture point, when all the fibres of component A are broken and the fibres of component B are partially stressed as per strain in the yarn. After this point the total load is borne by the fibres of the second component B. If component B is able to sustain the total load, the yarn reaches the second rupture point at a higher stress level. This is seen in cases 1 and 2. However, in the third case component B is not able to sustain the load and breaks immediately. These three cases are discussed here for prediction of yarn tenacity of blend yarns. Consider a blend yarn of tex $T$ having ‘$a$%’ of fibre A and ‘$b$%’ of fibres B, then

$$T_A = \frac{aT}{100} \quad \text{and} \quad T_B = \frac{bT}{100}$$
where $T_A$ is the tex of component $A$ and $T_B$ the tex of component $B$ and $T_A + T_B = T$.

When the strain reaches $X_1$, all fibres of component $A$ will be on the point of breaking and the total load $P_1$ supported by the composite yarn will be the sum of the loads $P_{1A}$ and $P_{1B}$ supported by the two components.

Where

$$P_{1A} = \frac{aTS_A}{100} \text{ and } P_{1B} = \frac{bTS'_B}{100}$$

if $S_1$ is the yarn tenacity at $X_1$, then

$$S_1 = \frac{P_1}{T} = \frac{1}{T} \left( \frac{aTS_A}{100} + \frac{bTS'_B}{100} \right)$$

$$= \frac{1}{100} (aS_A + bS'_B)$$

All fibres of component $A$ will break at this point. For the second rupture point at $X_2$ when all fibres of $B$ break, the total load $P_2$ supported by yarn would be the load supported by component $B$ that is given by:

$$P_2 = P_{2B} = \frac{bTS_B}{100}$$

If $S_2$ is the tenacity at $X_2$ then

$$S_2 = \frac{P_2}{T} = \frac{1}{T} \left( \frac{bTS_B}{100} \right) = \frac{bS_B}{100}$$

**Case 1:** Fibre $A$ has higher modulus but lower breaking stress and strain than fibre $B$. At the first rupture point when all fibres of component $A$ break, the stress shared by the component $B$ is lower than that of component $A$, i.e. $S_B > S_A > S'_B$. The resultant behaviour pattern of yarn tenacity shows that addition of the stronger and more extensible component $B$ in small proportion reduces the yarn tenacity. It is only after a certain minimum % of $B$ has been reached that the tenacity value becomes equal to that of 100% $A$ component. Beyond this value, the tenacity shows a continuous increase. Such behaviour is expected for polyester-viscose and polyester-cotton blend yarns as shown in Figure 8.10, where cotton and viscose are like fibre $A$, and polyester like fibre $B$. This behavioural pattern can, however, change drastically if the type of polyester and viscose happen to be of high tenacity.

**Case 2:** Fibre $A$ has lower modulus and lower breaking stress and strain than fibre $B$. In this situation, at the first break point when all the fibres of
component A break, the component B sharing a higher stress than that of component A, i.e. \( S_B > S'_B > S_A \). Such a blend would therefore show a continuous increase in tenacity of blend yarn with increase in proportion of component B, i.e. stronger fibre. There is no drop in tenacity with addition of component B even in very low proportions. Such behaviour is expected from blends of higher tenacity polyester and viscose rayon, and polyester and acetate.

**Case 3:** In this case, unlike cases 1 and 2, component B has lower breaking stress though higher breaking extension. At the first breaking point, the load shared by component B is also lower than that of component A, i.e. \( S_A > S_B > S'_B \). In such blends the addition of component B, the weaker one, reduces the yarn strength consistently and reaches the minimum value only when it is 100% component B. Such trends are observed when viscose or acetate is added to cotton. It must be mentioned here that the above analysis is applicable to single thread strength. The lea strength may show considerable deviation from these behavioural patterns because this test involves some other factors.

It should be understood that the assumption that all fibres of component A break at the first breaking point, is not quite true. Many studies have shown that some of these fibres break twice or even thrice. Such multiple fibre breakage means that a part of the broken fibre is still in integral part of the yarn and is capable of load sharing. Referring to Figure 8.9 it would mean that for the second rupture point the point \( S_B \) should not have been joined with the origin but slightly higher. The extent of such load sharing by the less extensible fibres is quite difficult to estimate. However, one can safely say that the actual values of tenacities would lie above the predicted ones in the middle region. In spite of these limitations, the simple analysis...
given here is of great help in understanding how the tenacity of fibres is realized in blend yarns, and gives a reasonably accurate prediction of yarn tenacity.

8.9.2 Breaking extension

Although yarn strength is considered to be the most important yarn property while selecting the right fibre for blending, the breaking extension of yarn should also be given due consideration. During shed formation in weaving, the breaking extension could significantly influence the weavability of the warp yarn. It is sometimes preferable to consider the work of rupture for this purpose.

In regard to this property, it has been found that an empirical approach is better suited. The change in breaking extension of a blend yarn with change in the blend ratio very much depends on the fibre extensibility of the blend components and the yarn structure. Figure 8.11 shows the breaking extension of some blend yarns at various blend levels. It can be seen that the shape of the curve is very different for the three blends. If the breaking extensions of the blend fibres are nearly equal as in the case of cotton and polynosic (C), there is practically no change in the breaking extension of the blend yarn. However, if the blend fibres have very different breaking extensions as in the case of cotton and polyester (A) or viscose and polyester (B) then there is an abrupt transition from a lower extension level to a higher extension level. In general, one could say that the breaking extension of a blend yarn lies in between the breaking extension of component yarns, that is the breaking extension of a polyester-cotton yarn.

![Graph showing effect of blend ratio on breaking extension of yarn](image-url)

**8.11 Effect of blend ratio on breaking extension of yarn:**
would lie in between the breaking extension of 100% cotton and 100% polyester yarns. However, it is rarely the arithmetic mean of the two. Unlike the yarn tenacity, it is extremely difficult to theoretically calculate the breaking extension of a blend yarn from the breaking extensions of the two components.

8.9.3 Yarn unevenness

Yarn unevenness is an important factor as far as yarn weavability and visual appearance of the fabric is considered. The main advantages of manmade fibres are the length uniformity and absence of short fibres. This results in improved drafting, as the fibre movement is under better control. This factor is of advantage in the blends of manmade fibres with cotton – especially carded ones – with a large length variation. Such length variation in 100% cotton mixing would give much higher irregularity during drafting (for example, as the viscose fibre content of appropriate length is increased in a cotton-viscose blend, the yarn irregularity goes down). For producing good quality yarns from polyester-cotton, it becomes essential, therefore, to improve the fibre length uniformity of the cotton component by combing it. The improvement in yarn evenness due to addition of manmade fibres is generally accompanied by a lower level of imperfections like neps, thick and thin places, and also of faults.

8.9.4 Abrasion resistance

It is supposed that abrasion resistance of the blend fibre is greatly affected by the component existing on the fibre surface. To find the composition at the fibre surface, the infrared reflection absorption spectrum was measured by the FTIR-ATR method. The result reveals that, with increase in PA6 content, its existence on the surface also increases.

To make high abrasion resistance fibres, polypropylene/polyamide 6 blend fibres of different blend ratios and viscosity ratios are being prepared by Takahashi et al. They finally concluded that the relationship between abrasion resistance and blend ratio of the blend fibres is found to be in a reverse-S-shaped curve as shown in Figure 8.12. The abrasion resistance increases with increase in PA6 content up to 20% of PA6 and then it follows a decreasing trend up to 60%. The abrasion resistance curve increases again as PA6 content approaches 100%. The result of the drawn fibre also shows a similar trend as can be observed from Figure 8.13.

In a blend fibre with the PA6 content range of 0–20%, the relatively rich PP component probably gives a sufficient wax effect to the surface of the blend fibre. In the PA6 content range of 0–20%, the breaking strength of the fibre itself increases, since the proportion of the PA6 component with
Polyesters and polyamides

high breaking strength increases as the PA6 content increases. The number of repetitions till abrasive rupture might have increased as the PA6 content increased. In the blend fibre whose PA6 content is more than 60%, another reason why the abrasion resistance increased with increasing the PA6 content might be the change of cross-sectional shape of the fibre.

8.10 Blended yarn structures

Kemp and Owen\textsuperscript{11} conducted experiments wherein blend yarns were extended to different levels of strain and then were taken apart. The
numbers of breaks of cotton fibres were then measured and the mean length of the cotton fibres was calculated. The breaks and the mean length are plotted for a 60/40 nylon-cotton blend in Figure 8.14 and here it is seen that a few fibres break at yarn extensions of 5% extension, but this number raises to 50 breaks at an 8% extension, to 100 breaks at a 10% extension, and to over 200 breaks at 15% extensions. The mean length of the fibre reduces correspondingly.

They also demonstrate that the resistance of the yarn to extension can be calculated from the weighted resistance of its individual components up to the level of about 6 or 7% extension, corresponding to the point where the lower extension fibre starts to break. Beyond this point it is necessary to determine the loss in load capacity due to the broken fibres, but also to take into account the further ability of the cotton fibre segments to contribute to the yarn tensile resistance. The assumption of the complete independence among the fibres of the yarn would lead one to predict that beyond 8% extension the yarn would have to depend entirely upon its nylon component, and there would be a significant drop in the load level at this point of the yarn extension test.

Machida\(^\text{12}\) predicted the stress–strain behaviour in a blended yarn and of the strength of yarns with varying blend ratios (Figure 8.15 (a)). His prediction was based upon the assumption of independent behaviour among the fibres of the yarn proposed by Hamburger.\(^9\) Figure 8.15 (b) shows the change in the height of the two rupture points which occur at the strains corresponding to the rupture strains of the individual fibre components. The yarn breaking strength is taken to be that rupture point which is higher for a given blend ratio.

\(8.14\) Mean fibre length and fibre breaks in a nylon-cotton blend.
Figure 8.16 shows the load-extension behavior of 4000 den model yarns made up of individual 70/34 nylon yarns and 100 s cotton yarns with a twist multiple of 3. The importance of fibre contribution to yarn strength adds significantly to the breaking energy of the yarn structure. Furthermore, it may be expected that when the blended yarn is introduced into a woven structure, the lateral force further enhances the contribution of cotton component to the breaking strength of the fabric at strain levels exceeding those of the cotton fibre. In Figure 8.17 there is clear indication that the actual breaking strength of the blended yarn falls above the predicted curve.
Furthermore, it is seen in Figure 8.17 that the tensile resistance of the blended yarn does not fall off rapidly after a 10% extension. The shaded portion in the stress–strain curves indicates the extent to which cotton, which must have ruptured at least once by now, continues to contribute to the resistance of the nylon-cotton blend. The question to which this study was addressed was how the broken lengths varied with strain and how they
varied with the structure of the yarn and their individual positions within that yarn. Note that this experimental work was done with model yarns in which the individual components, instead of being single fibres, were actually twisted yarns of nylon or of cotton.

The final question which remains, relates to the fibres which surround the cotton just at the moment that it breaks. As seen in Figure 8.18, the modification of the tensile distribution along the cotton fibre which occurs when breakage takes place somewhere between 2% elongation and 10% elongation must be accompanied by a corresponding modification of the tensile distribution along the adjacent closely packed blend fibre, in this case a polyester. Note in the right portion of Figure 8.18, that over the ruptured region of the cotton the neighbouring polyester fibre must suffer a much higher level of local stress, perhaps sufficient to cause its immediate rupture.

Clearly, one must now consider the relative strength level of cotton versus polyester so as to determine whether the rupture of the cotton will cause a catastrophic propagation of the breakdown to the surrounding blend fibres. The chances are that if the cotton is surrounded by other
cotton fibres in close contact, the break will propagate, since the surrounding cotton fibres would have already been loaded to a point very close to their breaking limit. Also, the additional load to be shared because of the breakdown of a neighbour would exceed their strength. Failure of the second and third and third and fourth cotton fibres in a location would not only increase the local load to be shared by adjacent fibres, but would also tend to reduce the pressure locally and thus cause increased slippage. One could consider an infinite number of variations of yarn breakdown, but it suffices to say that successful taking over of the load forfeited by an individual fibre by neighbouring fibres is desirable in an efficient blend yarn structure. And this would dictate the desirability of good dispersion of the fibres, that is, a minimum amount of clumping. Also, it would suggest the desirability of having the higher elongation fibre break at a tensile value considerably above that of the lower elongation fibre. Without this provision, it is likely that the blend yarn will fail catastrophically and the full tensile energy of the more extensible fibre components will not be realized.

8.11 Blending for speciality products

8.11.1 Core spun yarn

Core spun yarn has the strength and elongation of the central component and exhibits most of the other characteristics of the surface staple fibres. These yarns are widely used in a variety of applications; such as industrial clothing, conveyor belting, tentage and sewing threads. Core spun yarns can be manufactured by different methods like ring spinning, open end spinning, air jet spinning and friction spinning by the incorporation of certain attachments. In the case of staple fibre spun yarn, no single fibre is present parallel to the yarn axis due to the migration of an individual component. Migration occurs if all the components are delivered into the twisting zone at the same speed. In a true core yarn, therefore, the length of the component forming the core must be lower than the corresponding lengths of the covering component. The vital requirement for the ideal core yarn is that the covering fibres should enclose the core in a regular manner so that the core is completely buried at all points along the yarn length.14

Different methods for manufacturing core spun yarns

Core spun yarn can be manufactured by different methods like ring spinning, open end spinning, air jet spinning and friction spinning. In industry,
Core yarn is composed of two components, core and the sheath. The twisted central component of the core spun yarn lying parallel to the yarn axis, is termed as ‘core’. The core component may be mono- or multifilament. The outer twisted component covering the core is termed as ‘sheath’. Sheath fibres are either natural or manmade fibres. The properties of ring spun core yarn depend mainly on core-sheath ratio, and pre-tension applied to the core material. The core-sheath ratio depends on the core filament denier and yarn count. The major properties of ring spun core yarns (30s and 40s Ne) for different core-sheath ratios are given in Tables 8.1 and 8.2. The proportions of sheath (cotton) and core (polyester filament) used in these yarns are given in Table 8.3.

Yarn quality attributes are better with higher core component. To illustrate, when the core content is increased from 25 to 35%, yarn tenacity increases by 9% in 30s count and about 17% in 40s count. Breaking
Table 8.3 Proportion of sheath and core

<table>
<thead>
<tr>
<th>Count of core yarn</th>
<th>Core filament denier</th>
<th>Proportion of sheath/core</th>
</tr>
</thead>
<tbody>
<tr>
<td>30s</td>
<td>34</td>
<td>75/25</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>62/35</td>
</tr>
<tr>
<td>40s</td>
<td>34</td>
<td>78/22</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>68/32</td>
</tr>
</tbody>
</table>

elongation also improves by 1 to 2% (absolute values) and CV of tenacity reduces by 2 to 6% (absolute values). Evenness improves by 1 to 3% and imperfections decrease by about 35% when the core content is increased from 25 to 35%.

Core yarns are also manufactured in rotor spinning systems with a view to obtain higher productivity. For producing core yarn by open end spinning process, it is necessary to make some modifications in the conventional rotor spinning machinery. The modification of the rotor spinning process can be found in the literature. The advantages of producing core yarns on rotor spinning system, according to Lawrence and Chen are, (a) the rotor-spun core yarn is more likely to retain all the strength contributed by the core component and the full length of the core component, since the continuous filaments are not twisted during spinning and will therefore not suffer twist contraction, (b) an important consequence of this is that even metallic wires of high rigidities can be used as core, (c) the evenness of rotor-spun core yarns is better than that of equivalent ring spun yarn, and (d) the bulkiness of the rotor-spun core yarns is greater than that of the equivalent ring spun structure. Hence a reduction in cloth set will be required to obtain the same cover from rotor spun core yarns. Core yarns are also produced in DREF-2 and DREF-3 and air-jet spinning processes. The details may be found in the literature.

8.11.2 Blending in nonwoven and other products

Polyester polymers and copolymers are widely used in nonwovens in both staple fibre and spunbonded processes. The combined volume of polyester and polyolefin polymers used in the United States in nonwoven fabrics has been estimated to be more than 250 million kg annually. Blends of polyester fibres with wood pulp which are wet laid and then bonded by hydroentangling and subsequently treated with a repellent coating are widely used in disposable surgical gowns and drapes. Nylon fibres are used only
sparingly in the form of staple fibres and in a limited volume in spundbonded nonwovens. One of the largest uses for spundbonded nylon nonwovens is in the reinforcement of carpet pads and in fibreglass filters. The fabrics provide a low friction surface to carpet pads that facilitates the installation of carpets.

**Blends for filtration application**

Fine diameter fibres have always been of interest in filtration applications. A series of segmented synthetic splittable fibres has been developed that will split when processed through gentle carding systems such as cotton and semi-worsted cards. The fibres are produced from polyester and nylon 6 and are spun into a 16-segment 3 dpf × 51 mm crimped fibre (for cotton system) and other lengths and deniers can be produced for the semi-worsted system. The individual split fibre segments have effective diameters in the range of between 3–4 microns. The fibre is blended up to 30% of the fibre blend with other fibres. The fibre must be properly preblended and opened to maximize blend uniformity before carding. Worker and stripper loading must be considered when running this fibre because the split fibre has 16 times the number of fibres that must be processed through the system. A number of other polymer combinations are under development including polypropylene for electrostatic charging and blends of polymers for enhanced triboelectric properties. Figure 8.19 shows the fibre before carding.

**Figure 8.20** is a cross-sectional photomicrograph of the card-split fibre. The individual split segments can be clearly seen. Notice that the fibre

![Photomicrograph of card-splittable fibre before splitting](image-url)
segments are pie-shaped. This shape should be an advantage in filtration materials for lowering the clean pressure drop of materials produced from these fibres. Splittable synthetic fibres have been produced in short cut form with a water dispersible finish. Polymer blends include polypropylene/nylon 6 and polypropylene/polyester. The fibres have been produced as 16-segment 4.1 dpf $\times 0.24$” without crimp. When formed into handsheets with minimal opening processing, the physical properties of the handsheets match those of commercially available meltblown polypropylene materials as shown in Table 8.4. Split fibre segments range from 3–4 microns. Order of addition of furnish components will need to be addressed for each specific fibre blend produced.
These fibres have been successfully processed in fibre blends through a Rando Webber as high as 50% of the fibre blend. The fibre opened and split well in the machine and produced a uniform product with micro-denier fibre segments. When a card is used as part of the fibre opening system for the Rando (or the airlaid process is a Fehrer-type system), care should be taken to select a splittable fibre blend percentage that will successfully process through the card without jamming the worker and stripper sections.

The wetlaid shortcut version of the fibre has been successfully processed through a DanWeb Forming airlaid system up to 90% of the fibre furnish. The fibre was fed to the unit through a transport fan that opened and split the fibre into individual segments and formed it into an airlaid sheet. The resultant web was bonded with 10% binder fibre and was extremely soft to the touch. Blends with woodpulp and other fibres are expected to process well and can carry the splittable fibre through the hammermill section, if needed. Blend levels will require definition for each specific application.

Splittable synthetic fibres have been produced with cross-sections that will split when subjected to high-pressure water jets without splitting in the card. The fibres are processed as-spun into nonwoven products as standard textile fibres. The web is then passed through a hydroentanglement unit that splits the fibres and causes them to wrap around themselves and other fibres in the blend to enhance the tensile and toughness properties of the product.

**Blends with recycled fibres**

The textile industry provides ideal opportunities for developing technology for the use of recycled, reprocessed, or low-grade fibres to produce environmentally acceptable textile products. Many areas of textile processing for both garments and household goods produce waste materials that can be recycled into usable consumer products. More than 400 million tonnes of non-biodegradable polyester remnants are produced as waste each year. Such wastes provide good sources of materials for recycled products. In addition, most commercial cotton textile processes require that the fibres be scoured and bleached. Use of greige cotton in consumer products would not only reduce production costs, but would eliminate the discharge of chemicals needed in the scouring and bleaching process.

In a study, nonwoven blanket materials have been made from blends of recycled polyester fibres and greige cotton. Blending of cotton fibres into these nonwovens allowed application of finishes previously developed for cotton fibres. Nonwoven fabric structure was chosen because they are less expensive to produce than wovens, or knits, and because the short reprocessed polyester fibres could better be utilized in nonwoven processes.
Crosslinking finishes increased the stability of the fibres within the fabric, and also enhanced flame resistance. Finished fabrics withstood five laundry cycles without losing acceptable appearance or structural integrity. An antibacterial finish provided protection on unlaunched samples, but lost effectiveness as the laundering progressed. The fabrics were light-weight, had good hand, and excellent comfort and thermal qualities. These materials were expected to provide highly acceptable thermal coverings for refugee, medical, military, and recreational uses where a low cost, short-term use product is desired.

8.12 Summary

Polyester and polyamide fibres have interesting properties and these properties can be engineered to a specialized application during the process of manufacture. However, it is still not possible to obtain all the required properties for a particular application due to the inherent nature of these fibres. In these cases, it becomes essential to blend these fibres with other suitable fibres to enhance their properties. In this chapter, application of blending to three segments of applications, namely apparel end uses, core spun yarn applications and nonwoven and other applications are discussed in detail. Special attention has been provided on the evaluation of blending. Issues related to influence of fibre parameters on the blending behaviour as well as on structure and properties of the final yarn produced are discussed in length. In the case of blending of polyester and polyamide fibres for speciality products, core spun yarns, use of splittable fibres for making nonwoven filter products and blends using recycled fibres are discussed.

8.13 References

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Weaving technology for manufacturing high performance fabrics

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9.1 Principles of fabric formation

There are many ways of making fabrics from textile fibres. The most commonly used fabric forming methods are weaving, knitting, netting, tufting, braiding, lace making and nonwoven manufacturing. Among these methods, the most traditional and popular method of fabric manufacturing is weaving. Weaving is the interlacing of warp and weft yarns perpendicular to each other. There is a large variety of possibilities of interlacement between warp and weft yarns and the manner in which this is carried out determine the fabric structure. In woven structure the yarns are held in place due to inter-yarn friction. A prime requirement of textile fabric is that it should be flexible. Other requirements are very specific depending on the functionality desired in the fabric performance. The woven structures provide a combination of strength with flexibility. The flexibility at small strains is achieved by yarn crimp and by the freedom of yarn movement, whereas at high strains the threads take the load together giving high strength. In woven formation, great scope lies in choosing fibres with particular properties, arranging fibres in the yarn in several ways and organizing in multiple ways interlaced yarns within the fabric. This gives textile designer great freedom and variation for controlling and modifying the fabric. The yarn properties and the fabric structure together determine the properties of the fabric. The selection of fibre mix, yarn structure and fabric design predominantly depends on the end use application of the fabric.

Production of woven fabric starts with yarn preparation methods like winding, warping and sizing. During actual weaving process both warp and weft yarns are subjected to various complex stresses. The basic objective of yarn preparation methods is to prepare packages of desired size and build and also to make the yarns capable so that they can sustain all mechanical stresses during weaving. In fact the efficiency of weaving process and quality of fabric to a large extent depend on the quality of yarn
preparation. After satisfactory yarn preparation, weaving is carried out on a machine called a loom. Normally weaving machines are named after their weft insertion systems. Broadly weaving machines are classified as shuttle and shuttleless systems. Shuttle looms have been used for centuries and have now become obsolete in developed countries and partly being used in some industrialized countries for manufacturing of some special products. Shuttleless weaving machines emerged in the mid-twentieth century as potential weaving machines for mass production of high quality woven fabrics. Weft insertion mechanisms such as rapier, projectile, air jet and water jet are now being used for manufacturing various kinds of fabrics depending on the type of raw material, fabric structure and overall economics of the process.

9.2 Fundamentals of woven structure

9.2.1 Weave representation

The pattern in which the warp and weft yarns are interlaced is called weave. Practically there is unlimited number of weaves. This is in fact an obvious advantage that a weaving technologist can avail to produce unlimited woven structures differing in their appearance, properties and performance. A weave is symbolically represented by a weave diagram as shown in Figure 9.1(B) in which the columns represent the warp yarn and the rows represent the weft yarns. Each square represents the crossing of an end and a pick. A mark in a square indicates that the end is over the pick at the corresponding place in the fabric that is warp up. A blank square indicates that the pick is over the end that is weft up. Figure 9.1(A) shows

9.1 Plan (A), weave representation (B) and cross-sectional view along warp (D) and weft (C) of plain weave.
plain weave in plan view and cross-section along warp and weft are shown in (D) and (C).

### 9.2.2 Fundamental weave characteristics

The fundamental weaves are those in which every warp and every weft thread within the repeat overlaps or is overlapped only by one thread of the opposite system. Consequently, the number of warp threads in a repeat must be equal to the number of weft threads. Here, \( R_o = R_y = R \), where \( R \) = general weave repeat; \( R_o \) = repeat of warp; \( R_y \) = repeat of weft. There are three kinds of fundamental weaves: plain, twill and sateen weave. Each kind of fundamental weave is determined by two parameters, and each weave has its own values of parameters. Such parameters are: the repeat \( R \) of weave and the shift \( S \) of overlapping, or move. The shift is the distance between two neighbouring warp overlaps measured by the number of threads. There are two kinds of shifts: (a) Vertical shift (\( S_o \)) – the shift of two warp threads with respect to each other. This shift is often called the shift on warp where the count is made in vertical direction. (b) Horizontal shift (\( S_y \)) – the shift of two weft threads with respect to each other. This shift is often called the shift on weft, where the count is made in the horizontal direction.

The shift can be negative or positive. The vertical shift is positive, when the count is made upwards. The horizontal shift is positive, when the count is made from left to right, and negative when the count is made from right to left.

### 9.3 Basic weaves

There are three basic weaves such as plain, twill and satin weave. These are shown in Figure 9.2. However, innumerable derivatives can be developed from these three basic weaves; each of these fabrics has a different texture and properties.

![Basic weaves](image_url)
9.3.1 Plain weave

Plain weave has the simplest repeating unit of interlacement. It has a one up–one down interlacement of warp and weft yarns: as a result the fabric has the same texture on both sides. It also has the maximum possible frequency of interlacements and therefore has the maximum level of yarn crimp in the structure. Plain woven fabric has low modulus compared with other structures having less crimp. The weave gives an equal number of warp and weft overlaps in unit of weave. Plain weave fabrics are classified as balanced and unbalanced. In balanced fabrics the warp and weft counts are similar, and the ends and picks per centimetre are also similar. The yarn crimps are usually equal. Plain weave fabrics are widely used, much more than fabrics of any other weaves. In plain weave the values of parameters are the simplest: repeat, $R = 2$; shift, $S = 1$. There are only two threads with different interlacings within the repeat. Two heald shafts are sufficient to produce plain weave. When the number of ends per centimetre is large, four or six heald shafts are used with skip draft. Usually, tappet shedding motions are installed on looms for producing these fabrics. Plain weave is widely applied in various branches of the textile weaving industry. In trade, terms like tabby, calico, batiste are applied to plain weave fabrics.

9.3.2 Twill weave

The twill weave is produced in a stepwise progression of the warp yarn interlacing pattern which results in the appearance of a diagonal line in the fabric. The weave is widely used for ornamentation of the cloth. This design helps to achieve greater weight, closer setting and better draping characteristics of the fabric as compared with plain weave fabric produced from the same yarn. Twill lines are formed on both sides of the fabric; however, if warp float predominates on one side of the cloth, the weft float will predominate on the other side in the same proportion. A twill cannot be constructed upon two threads, but upon any number, more than two. The simplest twill contains three ends and three picks. In the repeat of fundamental twill the number of picks equals the number of ends. The twill is usually denoted by a fraction. The numerator of the fraction is equal to the number of warp overlaps and the denominator is equal to the number of weft overlaps within the repeat. The sum of the numerator and denominator of this fraction is the repeat of the twill. For instance, in $1/3$ twill, the number of warp overlaps within the repeat is 1, that of weft overlaps is 3, and the repeat is 4. When the shift is positive, the single overlaps form a diagonal which runs from left to right. This is usually called weft face right-hand twill. But when the shift is negative ($S_y = -1$) the diagonal...
runs from right to left and the weave is called left-hand twill. In the other weave, i.e. 3/1 twill, the repeat of the weave is 4. On each thread within the repeat, there are three warp overlaps and one weft overlap. This is a warp-face twill. This weave has a warp effect in contrast to twill 1/3 which has a weft effect, since the proportion of warp overlaps to weft overlaps within the repeat is 1:3. While designing the fabric with a warp effect, it is preferable to select the density of warp threads higher than that of weft threads, and vice versa. It is common practice to use straight draft for producing twill fabrics.

9.3.3 Sateen weave

In sateen weave, one yarn has a long float over several of the other yarns on one side of the fabric. This produces fabric with a high degree of smoothness, lustre and without any prominence of weave feature. Sateen weave enables one to produce fabrics of close thread packing and heavy construction. In pure sateen weaves the surface of the cloth consists almost entirely of weft floats, as in the repeat of weave each thread of weft passes over all and under one thread of warp. In addition, the interlacing points are arranged so as to allow the floating threads to slip and cover the binding point of one thread by the float of another, which results in the production of fabric with a maximum degree of smoothness and lustre and without any prominent weave features.

The parameters of sateen weaves are: $R > 5; 1 < S < (R - 1)$. Besides, for the construction of the regular sateen the shift ($S$) and repeat ($R$) must be expressed by prime numbers, i.e. they must have a common divisor but unity. Sateen weave is denoted by a fraction. The numerator of fraction is equal to the repeat of the weave. The denominator is equal to the shift ($S_y$) of overlaps. The warp face fabric is called satin. Satin usually can be constructed using the vertical shift ($S_o$). Satin has the warp effect, and the density of warp is much higher than the density of weft.

9.4 Theoretical considerations in woven structure

The use of different weave varies the ability of component threads to move relative to one another; as a result, mechanical properties like shear characteristics and drapeability of fabric change significantly. Therefore, the design of a fabric to meet the requirements of a particular end use is a complicated engineering problem. Theoretically, it is possible to design a fabric structure to achieve any desired characteristic, but in actual practice it is not so easy because of inherent non-linearity and complex relationship between structure and properties of textile materials along with their viscoelastic behaviour. The factors associated with fabric design include fibre
Polyesters and polyamides

empirical relationships between some fabric parameters and the fabric properties. There are also some established theoretical relationships based on fabric geometry and simple mechanics using first principles mostly for plain woven fabrics. The formal structure of a woven fabric is defined by weave, thread density, crimp and yarn count. The interrelation between fabric parameters can be obtained by considering a geometrical model of the fabric. This model is not merely an exercise in mathematics but useful in determining the entire structure of a fabric from a few values given in technological terms. It also establishes a base for calculating various changes in fabric geometry when the fabric is subjected to known extensions in a given direction or known compressions or to complete swelling in aqueous medium. Mathematical deductions obtained from simple geometrical form and various phenomena in fabrics. In this chapter a basic Pierce’s model for plain weave is shown in Figure 9.3. It represents a unit cell interlacement of a plain woven fabric in which the yarns are considered inextensible and flexible. The yarns have circular cross-sections and consist of straight and round segments. From the two-dimensional unit cell of a plain woven fabric, geometrical parameters such as thread-spacing, weave angle, crimp and fabric thickness are related by a set of equations. The symbols used to denote these parameters are listed below.

- $d$, diameter of thread
- $p$, thread spacing
- $h$, maximum displacement of thread axis normal to plane of cloth (crimp height)
- $\theta$, angle of thread axis to plane of cloth (weave angle in radians)
- $l$, length of thread axis between the planes through the axes of consecutive cross-threads (modular length)
Weaving technology for manufacturing high performance fabrics

• $c$, crimp (fractional)
• $D = d_1 + d_2$

Suffixes 1 and 2 to the above parameters represent warp and weft threads respectively. In the above figure projection of yarn axis parallel and normal to the cloth plane gives the following equations:

$$c_1 = \frac{l_1}{p_2} - 1 \quad [9.1]$$

$$p_2 = (l_1 - D\theta_1)\cos\theta_1 + D\sin\theta_1 \quad [9.2]$$

$$h_1 = (l_1 - D\theta_1)\sin\theta_1 + D(1 - \cos\theta_1) \quad [9.3]$$

Three similar equations are obtained for the weft direction by interchanging suffix from 1 to 2 or vice-versa.

$$c_2 = \frac{l_2}{p_1} - 1 \quad [9.4]$$

$$p_1 = (l_2 - D\theta_2)\cos\theta_2 + D\sin\theta_2 \quad [9.5]$$

$$h_2 = (l_2 - D\theta_2)\sin\theta_2 + D(1 - \cos\theta_2) \quad [9.6]$$

$$d_1 + d_2 = h_1 + h_2 = D \quad [9.7]$$

In all there are seven equations connecting 11 variables. If any four are known then the equations can be solved and the rest can be determined. These fundamental relationships can be used by fabric designers to decide the specification of fabrics according to their end use.

### 9.5 High performance fabric

Industrial textiles are specially designed and engineered structures. Unlike ordinary traditional clothing and furnishing fabrics, industrial textiles are generally used for various high performance or heavy duty applications. Synthetic fibres offer high strength, elasticity, uniformity, chemical resistance, flame resistance and abrasion resistance. In fact flexibility, elasticity and strength are simultaneously found in some of the textile materials that no other industrial product can provide. The development of exceptionally strong high performance fibres widened the area of applications of industrial textiles. High performance fibres offer special properties due to the demands of the respective application. These demands cover properties such as high strength, high elongation, high modulus and resistance to heat, chemical and environmental attack. Glass fibre, carbon fibre and aramid fibre are among the best known high performance fibres used in production of various speciality applications under high performance fabrics.
Carbon fibres containing high strength-to-weight and high stiffness-to-weight ratio enabled production of fabrics which are thermally and electrically conductive and exhibit excellent fatigue resistance. Fabrics woven from aramid and other high performance yarns are used for ballistic protection both for civilian and military personnel. These fabrics produced from Kevlar offer unmatched resistance to bullet, fragments, cuts, stabs and abrasion while offering light weight, high strength and comfort. Blast resistant fabrics produced from high performance para-aramid fibre display excellent dimensional stability over a wide range of temperature for a prolonged period. There are several soft as well as hard body armour products now being developed using woven fabrics from Kevlar. The applications are ballistic vest and helmet, racing helmet, ballistic blanket and electronic housing protection.

9.6 Yarn preparation for high quality fabric

To ensure high productivity and economic efficiency in a weaving mill, good warp preparation is indispensable. Faultless warps are an essential precondition for utilizing assets in the best possible way and achieving the fabric quality demanded by the market.

Yarn preparation for weaving depends on fundamental understanding of the yarn to be processed and the knowledge of stresses to which the yarn would be subjected to during weaving. Modern weaving machines have placed increased demands on warp preparation due to faster weft insertion and use of different weft insertion devices. High efficiency is crucial to ensuring economic production. It is unrealistic to assume that a high quality warp can be produced with an inferior quality yarn. On the other hand, it is possible to produce an inferior quality warp from a high quality yarn, and to avoid this, a number of important points should be kept in mind when making warps. For example, a knot in the weaving room can cause a warp stoppage and possibly a fault in the fabric. For this reason a correct weaver’s knot should be used for every broken end. Nowadays the use of mobile splicers can be recommended in weaving to mend broken ends which may cause faults in the fabric. Yarn preparation during warping and sizing is crucial for both loom shed efficiency and fabric quality.

9.6.1 Warping

In warping, hundreds of yarns drawn from individual packages lie parallel to one another, and are wound over a drum. During this process a preliminary assessment of yarn quality can be made on the basis of stoppage records or analysis. This is especially important with bought-in yarns which
are utilized without being inspected on receipt. For good warps and economy in the subsequent processes the stoppage value of the order of 5 per 107 metres of ring spun spliced warp yarn is considered as satisfactory in the warping room. The following are some of the measures that can be taken to minimize stoppages:

- prevention of fly by regular cleaning – no weaving machine stoppage due to oversized fly;
- creeling of cylindrical packages with the same unwinding direction – uniform unwinding direction avoids collision of balloons;
- avoidance of lost warp ends – no stoppages at the sizing machine. This is accomplished through good training of operators and regular maintenance of the warping machine, e.g. correct adjustment of the stopping distance and correctly functioning stop motions, etc.

When winding the yarn sheet onto the warp beam, care must be taken to ensure uniform distribution of the yarns. The beam must not have any excessively high or sloping edges; it must be completely cylindrical. For warping of synthetic yarns various measures like use of chemicals, ionization of air or humidification of air can be used to eliminate static charge. Special tension rollers need to be used between creel and warper while warping heavy industrial yarns to maintain desired thread tension.

### 9.6.2 Sizing

During sizing the warp yarn is temporarily coated with a protective layer of adhesive. It gives the warp yarn the resistance necessary to sustain weaving stresses on the loom. If the sizing machine is correctly set and the correct sizing agent has been chosen, the strength of the yarn is improved and its hairiness reduced. At the same time the sizing process also reduces yarn extensibility, which has a major impact on the warp breakage during weaving. In the case of ring and OE yarns, this reduction should not be more than 25% of the breaking extension of the parent yarn. A modern sizing installation for spun yarns should be equipped with pre-wetting attachment, two size boxes depending on the yarn cover factor of the squeeze rolls, wet splitting arrangement with separate drying, tension regulation system, regulation of squeezing pressure, regulation of residual moisture, measurement and regulation of size application, constant winding tension and after waxing device. Pre-wetting results in 25–30% saving of sizing agent. With this system, size application is restricted to the yarn surface and it offers uniform coating and better size–fibre interaction, which have a positive impact on the weaving process. Compared to conventionally sized warps, pre-wetted warps exhibit better weavability and
reduced warp stoppages. An appropriate choice of sizing agent and optimization of the size recipe and process parameters have a major impact on the weaving performance of the warp yarns. The sizing machine should not be stopped at all during running. All possible measures should be taken to run the machine at uniform speed. Stretch of warp should be minimized and controlled accurately to preserve residual elongation in the yarn to sustain weaving stress. A balance between size penetration, size coating, add-on, yarn hairiness, residual yarn elongation and yarn abrasion resistance is essential to achieve high weaving efficiency. For instance, warp beams are often not filled to their full capacity. Over the years, additional warp changes cause extra costs which can be avoided by appropriate planning and working methods. The use of the right transport apparatus, knotting machines, drawing-in equipment, etc. can have a considerable impact on weaving room output.

Filament warp yarns can be made weavable by one of the three different means, i.e. sizing, twisting and intermingling. In all the three alternatives, cohesion between the neighbouring filaments of the multi-filament yarn is generated which holds the fibres together and reduces separation and filamentation during weaving. In the case of twisting as well as intermingling, the yarn’s physical structure changes, whereas sizing is the only process which improves the weavability of the yarn without disturbing the structure and surface characteristics of the warp yarns. Both twisting and intermingling deteriorate some of the mechanical properties of the yarn particularly when the filaments are very stiff and brittle. However the choice of a particular process is largely dependent on the end use application, process cost and quality of the fabric to be produced. For sizing filament yarn single end sizing principle is most suitable to prevent yarn flattening and filament sticking. The important task in filament sizing is control and monitoring of warp tension as these yarns are highly susceptible to extension.

9.7 Weaving systems

Weaving machines are known according to the weft insertion systems being employed on the loom. Weft insertion with a shuttle is the oldest weaving system in which weft is inserted by a shuttle that traverses back and forth across the loom width. Since the weight of the shuttle is several thousand times greater than the weight of the weft to be inserted in a pick, it is considered as an inefficient process and has become obsolete. Alternatively, other weft insertion systems have been developed and they are running successfully in industry. The most popular shuttleless weaving machines are air jet, water jet, rapier and projectile machine.
9.7.1 Air jet weaving

In air jet weaving system, the weft yarn is inserted into the warp shed by a blast of compressed air coming out of a nozzle. Normally the air velocity in tandem and main nozzles exceeds sonic velocity and provides high initial acceleration. Subsequently as the air expands freely, relay nozzles are used to maintain high air velocity across the shed. A profiled reed is also used to guide the air stream and to separate the weft yarn from the warp. The flow of air during air jet insertion is unsteady, turbulent and either incompressible or compressible depending on the velocity. The transfer of weft in air jet insertion takes place due to the propelling force generated by friction between the air and the yarn surface. The amount of propelling force generated during weft insertion is given by:

\[ F = 0.5C_f \rho (U - V)^2 \pi d l \]

where \( C_f \) = skin friction coefficient, \( \rho \) = air density, \( U \) = air velocity, \( V \) = weft velocity, \( d \) = yarn diameter, \( l \) = yarn length in air.

The propelling force is directly proportional to the square of the relative velocity between the air and weft yarn. This force also increases with the increase of yarn diameter. Skin friction coefficient depends on surface characteristics of the yarn. The air jet weaving machine offers the highest weft insertion rate among all shuttleless weaving machines. Because of high productivity, these machines are mostly used for the mass production of standard fabrics such as sheeting, denim, terry towel, glass fabrics, institutional clothing material and tyre cord. The air jet weaving system can process natural, synthetic, spun and filament yarn for production of a wide range of styles. Textured yarns in the weft direction are preferred as they generate very high propelling force. Monofilament yarns are not suitable for air jet weaving as they have smooth surface and lack enough friction between air and yarn for propagation in the shed. In general the suitability of a yarn for pneumatic insertion depends on the count, twist and structure of the yarn and it is independent of the fibre material. Since the machine works at very high speed the quality of yarn and the slashing quality have to be of very high standard, failing which there would be significant loss in production and efficiency. Air jet weaving machines are also used for manufacturing of tent fabrics, airbags, parachutes, etc.

9.7.2 Water jet weaving

In water jet weaving, the weft yarn is drawn through the warp shed by means of a highly pressurized stream of water. The tractive force is provided by the relative velocity between filling yarn and water jet coming out of a nozzle. This drag force can be affected by the viscosity of the water...
and roughness and length of the filling yarn; higher viscosity causes higher forces. The drag force accelerating the weft can be written as:

$$D_f = 0.5 \rho_w C_f \pi \frac{d^2}{L} V_r^2$$

where $\rho_w =$ water density, $C_f =$ drag coefficient, $d =$ diameter of weft, $L =$ length of weft and $V_r =$ velocity of jet relative to weft. The basic principle of weft insertion with water jet is similar to that of air jet as both systems use a fluid to carry the weft yarn. However, one prerequisite for water jet weaving is that the yarn has to be wettable so that it can develop tractive force. In a water jet system, the propulsive zone is elongated as the water jet is more coherent due to viscosity and surface tension of water. Since the wet moving element is heavier, the probability of weft yarn to entangle with the warp line is very meagre. The weaving system is advantageous in terms of energy requirement, noise level and the jet is broken into droplets which create very little turbulence to disturb the weft yarn. However, the droplets spread in the shed and wet most part of the warp which restricts the use of this machine for the warp sized with water soluble adhesives. Therefore, water jet weaving is normally restricted to filament weaving. Modern water jet looms have a speed of about 1500 ppm, the maximum reed width can go up to 3 m and weft insertion rate can go as high as 2000 mpm. Light weight synthetic apparel cloths, tent/tarpaulin fabrics, inter-linings and canopy fabrics and high density air bag fabrics are made on these looms.

### 9.7.3 Rapier weaving

In this type of weaving a flexible or rigid solid element called a rapier is used to insert the weft yarn across the shed. The rapier head picks up the weft yarn and carries it across the shed. After reaching the destination the rapier head returns empty to pick up the next filling yarn, which completes the cycle. The conventional grippers are redesigned to ensure better clamping of the yarn and prevent rubbing of warp yarns. A rapier machine could be of single or double rapier type. The single rapier is normally a rigid rapier which carries the weft yarn from one end, passes it across the weaving machine and returns back empty. Therefore half the traverse time is wasted and loom width can be maximum up to the length of the rapier which requires more space per unit reed space. However, this has only one advantage that it can handle the weft yarn which is otherwise difficult to control as there is no yarn transfer from rapier to rapier. In the case of a double rapier system, the transfer of yarn takes place at the centre of the loom resulting in only half of the rapier movement being used for weft insertion. A double rapier machine can be either rigid or flexible which makes the difference in space requirement.
Rapier weaving machines are known for their versatility. These looms can weave very light fabrics of 20 gsm to the heaviest fabric of the order of 5000 gsm. The gripper heads can take a wide range of yarn count ranging from 5 to 1000 tex. It can manage successfully up to 16 different weft yarns enabling the weaver to weave any kind of fancy fabric along with an electronic jacquard. Apart from fancy and furnishing fabrics, rapier machine can easily handle filament yarns for manufacturing technical textiles. This is because of the fact that the rapier system of weft insertion exerts minimum stress on the weft yarn. The stress in weft yarn increases with increase in machine speed and it also depends on the elastic modulus of the yarn. Contrary to other shuttleless weaving systems, the speed of the rapier weaving machine is limited far more by the properties of the filling yarn than those of the warp. In the latest machines rapiers are made of composite materials and the rapier guide is eliminated. These machines are used for making automotive fabrics, fabrics for aircraft defence industries, heavy filter cloth and lightweight sports wear.

9.7.4 Projectile weaving

Projectile weaving machines use a projectile or a gripper miniature shuttle to insert the filling yarn across the machine. The gripper grips the end of the weft yarn presented to it and is projected across the warp shed. The projectile does not have to carry the weft package with it and it is therefore much lighter compared with the shuttle. The force needed to accelerate the projectile is also less and the picking mechanism is obviously lighter. However the mass of the projectile is heavy enough to be unaffected by minor obstructions in the warp shed. Since the mass of the projectile is much lower than that of the conventional shuttle, the speed can be increased substantially. The acceleration of the gripper can be increased by a factor of about 7 which offers advantage in terms of productivity and space. This unique weft insertion system uses the torsion bar picking system in which the strain energy is stored by twisting the torsion bar prior to the picking and it is released during the acceleration of the projectile by a toggle action. The projectile glides through the shed in a rake shaped guide. On its arrival at the other end, it is received in a specially designed unit and then conveyed back to its original position. The size of the projectile is small both longitudinally and in transverse direction compared with the shuttle size. The time required for it to pass through a given point is less and also the warp shed need not open too much for easy departure of the gripper. Therefore, a tighter loom timing can be accommodated to take the advantage of increased loom speed or larger fabric width. Projectile weaving practically allows the use of any type of yarn: cotton, wool, mono or multifilament, polypropylene ribbon and even hard fibers like jute and linen.
It is widely used for manufacturing cotton felts, agro-textiles, geotextiles, conveyor belts, cinema theatre screens, tarpaulins, paper machines, clothing and tyre cord.

9.7.5 Multi-phase weaving

The multiphase weaving machine is one in which several phases of the working cycle take place at any instant such that several filling yarns can be inserted simultaneously. In this more than one shed is formed at a time. The multiphase weaving system at present can weave 190 cm width with 69 m of fabric per hour. The weft is inserted continuously without interruption with an even pull off speed of around 20–25 m/s thus the stress on yarn is reduced. The system is more suitable for harsher bast fibres and cottons but unsuitable for weaving of continuous filament yarns. They have the added problem of stopping the loom in the event of weft break in any of the sheds. As four sheds operate simultaneously the system is very complicated with many small parts operating together. The machine can weave plain, 2/1, 3/1 and 2/2 weaves; and warp density is limited to a maximum of 45 ends per cm. Because of these drawbacks these looms are not commercially popular. It weaves fabrics of limited variety but at a substantially faster speed (about 5500 mpm) than any other conventional high-speed weaving machine.

9.7.6 Circular weaving

In circular weaving machines the warp is circular and there are continuously circulating shuttles running around the periphery in a wave or ripple shed. Circular fabrics are tubular fabrics of varying diameter without a lateral fold or edge. Tubular fabrics woven on flat weaving machines have a folded edge as they are joined at selvedges. In this the shuttles require a continuous motion across the shed and cannot leave the shed. On circular weaving machines a ripple shed is formed in which the warp is divided into segments which forms a shed with small heddle frames. An automatic shuttle change mechanism is used to remove the empty shuttle at a particular spot in the shed and a reserve shuttle is inserted. Drop wires pressing the filling yarns towards the fell of the fabric perform the beat-up operation. The machine is used for production of tubular fabrics for special applications such as woven sacks, tubes, medical textiles, etc.

9.7.7 Three dimensional weaving and composites

Three dimensional (3D) woven fabrics are fabricated by modifying the conventional weaving mechanisms. Harnesses with multi-eye heddles are
used to arrange the warps into three sections in plane form for weaving convenience. Mainframe and flanges are interlaced by a set of warps moving to and fro as a joint. Weft passes through the clear warp sheds separated by multi-eye heddles to form the 3D woven fabrics in plane form. The differential feeding length between the warp yarns gives rise to extra friction, and therefore hairiness may occur. In order to reduce this friction the warps are passed through the tensioner and weight with ceramic eyes individually between the creel and weaving loom. The thickness of the central portion of the flattened fabrics is different from the side portions. Therefore the cloth roller cannot be used to take up the flattened fabrics. The fabric is clipped and pulled by a pair of rollers set in front of the loom as a take-up device.

The technology is used for the production of only speciality industrial fabrics such as for making preforms for construction, automotive, ballistic and various industrial uses; for marine applications such as carbon fibre preforms for high performance powerboats; in medical technology (artificial veins, arteries, orthopaedic tubes); lightweight construction (reinforced section in automotive engineering and aeronautics); pipeline construction; in sports like shinguards for soccer, protective headgear for skydiving, high speed water sports, etc. Preforms made by 3D weaving provide several important advantages in composite fabrication. The most important advantage of this material is seen in manufacturing thick composites, owing to a significantly reduced labour time, when multiple layers of 2D fabric plies are replaced by one or a few 3D plies to achieve the required thickness in a composite structure. It is obvious to expect that the processing advantages of thick 3D woven preforms come at the expense of reduced conformability. In fact 3D preforms appear to be better than the most conformable 2D fabrics. The flexural, tensile and compressive stiffness and strength are better in laminates made from 3D preforms than those made from comparable 2D woven or even knitted fabrics. This is mainly due to the absence of in-plane crimp of yarns in the materials.

Woven composites have proved ideal for security applications, where a high level of protection is expected from the lightest possible components. New areas of application are under development. In transport applications and vehicle construction, in particular, composites have major advantages. The lower weight results in a higher payload with lower fuel consumption. To meet the high standards required, fabrics woven from yarns containing appropriate special fibres and with sufficiently strong constructions are needed. Special criteria for the composite can be fulfilled with specific materials, such as appropriate synthetic fibres. Materials of this kind are usually designed on an interdisciplinary basis by specialists in various fields. State-of-the-art computer analysis with the finite-element method
allows the properties of the fabric to be reliably predicted. Depending on the application, the component is built with different materials. In most cases, the fabric inlays perform important functions. The combination of different raw materials and fabrics with the matrices gives the desired properties.

The fibres used in the production of fabrics for composite materials are mainly inorganic and modified synthetic fibres, such as glass and carbon fibres, para-aramids, high-strength PE fibres, etc. For optimal take-up of forces by the fibres, the yarns are mainly of the untwisted multifilament type. However, spun yarns with a matrix in the yarn are now also being used. The design of the fabric is also dictated by the application. They are not standard fabrics produced in long production runs with unchanged settings. Exact reproducibility must therefore be assured.

## 9.7.8 Multi-directional weaving

Multi-directional weaving enables placement of fibres and yarns in various directions to obtain the required weave architecture and physical and mechanical properties. Fibre type, direction, spacing and volume fraction are key variables to achieve specified strength, modulus, density, electrical and thermal and many other transmission characteristics of the fabrics. Applications requiring multi-directional woven structures usually are those where extremes in temperature and highly stressed states are encountered. Various high strength, high modulus and high temperature resistant fibrous materials in tow or yarn forms are used in weaving. Fibres and filaments used to develop specialty weaving architectures include quartz, zirconia, silicon carbide, carbon, graphite, tungsten and impregnated yarns. Various fibres can also be woven into the same preform (hybrid weaves). Composite preforms can be woven with fibres oriented in three directions, as with orthogonal (mutually perpendicular) or polar (cylindrical) constructions. In addition, fibres may be oriented in 4-, 5-, 7- or 11- directions, which are referred to as multi-directional woven constructions. These woven structures are used either as preforms or as impregnated components for fabrication of fibre reinforced composites.

## 9.7.9 Jacquard

Grosse Webereimaschinen GmbH, Germany, and Stäubli developed the UNISHED and the UNIVAL 100 jacquard technology with a different concept. Although the principle of shed formation of these two machines is different, but they have a common goal to reduce the number of jacquard engine parts. The shed formation in the UNISHED is achieved using leaf springs. Each leaf spring is connected to a heddle that controls one warp
end. The leaf springs, which are controlled by actuators, control the bottom shed as well as the top shed. The configuration of the jacquard head and the individual control of each heddle or warp end allow the heddles to be set vertically. These settings eliminate the need for harness cords, magnets, hooks, pulleys, springs and the gantry. This results in lower building and air-conditioning costs. The jacquard head is mounted directly on the side frames of the weaving machine, thus making quick style change (QSC) possible in jacquard weaving, as it is easy to exchange the entire jacquard head, including the heddles.

9.8 Production of some speciality fabrics

Woven fabrics with special features are now used in several applications. With advancement in manufacturing technology and invention of new high performance fibres, the use of industrial fabrics is increasing in agriculture, civil engineering, protection and safety, automotive industry and transportation, storage and packaging, medical and ecological sectors, sports and recreation, electronics and instrumentation, etc. Some of the technical textiles have replaced conventional textile materials and interestingly many technical textiles have replaced building materials and metals mainly because of their low weight, high tenacity and resistance to chemicals and corrosion. The production of industrial textiles requires high tech weaving machines such as projectile, rapier and air jet weaving machines with auxiliary attachments for specific purposes. Only a few examples are discussed in this chapter because of space constraints.

9.8.1 Airbag fabric

Airbag is one of the most important safety devices used in vehicles such as aircraft, motorcars and para-gliders. It is installed inside the vehicle in front of a seated occupant to provide protection against injury arising due to collision with the fixed portion of the vehicle body during an accident. Therefore the airbag must inflate and provide a cushion for the occupant immediately after the collision. The material which inflates and provides the cushion is typically a woven textile fabric. For better performance of the airbag, the fabric has to meet certain performance requirements such as low air permeability, high bursting strength, high tear resistance, good packability and good edgecombing properties. In order to achieve these multifarious characteristics the fabric must be designed carefully from the selection of the fibre material till final finishing. The main requirements in airbag fibre materials are high strength, heat stability, good ageing characteristics, energy absorption, coating adhesion and functionality inextreme hot and cold conditions. The most widely used yarn in airbag manufacture
is nylon 66 yarn at a denier ranging from 420 to 840. Nylon 6, nylon 46 and polyester multifilament yarns are also used.

Airbags are made of compact plain woven fabric. Normally a thread density of about 18–21 threads per cm are chosen for both warp and weft direction to weave the fabric. Initially conventional rapier looms were used to manufacture the fabric. Subsequently high production water jet weaving machines are also being used successfully. The basic requirement of a weaving machine to produce airbag fabric is its ability to weave high thread density so that a compact structure can be made to provide low air permeability. In some situations to compensate for the lower weave density achievable on available weaving machines, weaving can be performed using yarns having high breaking tenacities so as to provide improved strength in the final fabric despite the lighter weave construction. Multifilament yarns for warp may require sizing with some synthetic binders such as polyvinyl alcohol, polystyrene and polyacetates to enhance the mechanical integrity of the flat filaments during weaving. Although these sizing compounds are typically effective in enhancing the mechanical integrity of the high tenacity yarns, such sizing also tends to enclose yarn oils which may not be compatible with polymeric compounds used for coating the fabric prior to its formation into an airbag structure. Therefore it is necessary to eliminate the sizing compound as well as the enclosed yarn oils by the scouring and drying of the fabric prior to making any coating operation.

9.8.2 Tyre cord

The tyre is a complex technical component and performs a variety of functions. It has to cushion, dampen, assure good directional stability, and provide long-term service. The most important characteristic of a tyre is that it should be capable of transmitting strong longitudinal and lateral forces in order to assure optimal and reliable road holding qualities. The pneumatic tyre is a rubber/textile composite. Cotton has been used as reinforcement in tyres for a long time. Continuous filament rayon began to be used for tyres in the 1930s. Research subsequently led to high tenacity variants and much needed improvements in fibre rubber adhesion. Normally the fibres used for tyre cords are high tenacity (above 6 g/den) filament yarns of nylon 6 and nylon 66, polyester and viscose, etc. While HT viscose and nylon were preferred earlier, they have now been replaced by polyester fibre which has lower creep under load and better heat resistance. Cords comprising the breaker or belt layers, however, use high modulus fibres, e.g. para-aramid (Kevlar), glass or steel. However, it is now realized that viscose is better than nylon in a high-speed impact. Polyester loses advantage of strength above 150°C. At 160°C, polyester shrinks by 2% in
length, nylon by –4% and viscose by just 0.1%, Viscose is again the fibre of choice in radial-ply tyre casing. In the technical information literature of Michelin tyres, relative properties of tyre cord materials are compared, reporting that the worldwide consumption of textile fibres in tyres is more than 744 million tonnes and fibre use share is 57% Kevlar, 24% polyester, 19% nylon.

Tyre cord is a warp dominant textile structure. Normally the warps, with a density of 6–13 threads per cm, are kept in parallel and only one weft thread per cm is used. Specially twisted, high strength filament yarns of polyester, polyamide, aramide or rayon are used for the warp. The weft material is usually a cotton or core spun yarn. Due to the low weft density the productivity of a machine weaving tyre cord is extremely high. Weaving machines with different weft insertion systems are used, depending on the type of fabric. Normally for tyre cord production, machines with working widths of 190 and 220 cm are used. To achieve long yardages all weaving is done from a modern creel unit and the fabric is wound into large rolls on a digressive batch winder. Instead of being fed from beams, the warp is fed directly from bobbins placed on creels which are accounted for by the fact that the number of warp yarns in the cord fabric is comparatively small and the warp is not sized. Instead of being wound onto the take-up roller the finished cord fabric is wound onto a special out-built stand mounted outside the loom area. This is done for the reason that the cord fabric is very thick, and the rolls of fabric are very large, thus necessitating greater overall dimensions of the loom and rendering its maintenance very difficult. Some mechanisms on looms are simplified; instead of temples used in standard looms to control the normal fabric width, the cord looms are provided with lamellar limiters. There are two healds in the shedding motion. The main plies of the carcass are manufactured from a dense cord fabric (distance between warp yarns 0.24–0.39 mm). Intermediate plies of looser cord fabric with a distance between warp yarns of 0.52–0.61 mm are laid between the main plies and the breaker. The two upper layers of the tyre carcass are called the breaker; it is intended for safely joining the elastic carcass with the stiff thread. The breaker plies are made from a cord fabric with a low count warp, the distance between warp threads being 1.29–1.36 mm. The plies of the breaker cord fabric are laid over the other at an angle of 45°.

Tyre cord fabric is a skeletal structure which holds the uniformed rubber mass of the tyre. The essential properties of this fabric are high tensile strength, less extensibility and adhesion compatibility with rubber and rubber chemicals. In tyre manufacturing technology, the tyre cord warp sheet (fabric) plays a major role. The overall strength and ability to bear load by the tyre depends strongly on the nature of the cord. The tyre cord should have high tenacity (usually above 6 g/den), high cut-through
Polyesters and polyamides

resistance, good impact resistance, good flex fatigue resistance, good thermal stability, capability to strongly bond with the rubber, low elongation and low moisture regain, high grip and rolling resistance, shock resistance, dimensional stability and elastic recovery.

9.8.3 High performance light weight foldable stretcher fabric

The stretcher forms an essential item in the inventory of any army. Based on extensive research and the design requirements of the stretcher, the fabric has been woven using nylon 66 yarn; Denier of yarn: 500; No. of plies in yarn: Single; No. of filaments per ply: 140; Tenacity of yarn: 40 cN/tex; Texturized yarn. This fabric has got excellent mechanical properties and also offers resistance to bacteria, mildew growth, fungus, water (especially sea water), and corrosion. It is 60 per cent lighter than canvas fabric presently being used. This fabric has numerous other defence applications. It has all-weather capabilities, takes less drying time and offers ruggedness, strength, endurance, excellent abrasion resistance, good tear and puncture resistance, extreme fade resistance, etc. The fabric can be used for making combat uniforms, high altitude and winter clothing, tents, hovercraft, uppers of jungle boots, harnesses, covers, luggage, web equipment, ground sheets and parachutes. Areal density: 272 gsm; breaking load (5 cm × 20 cm strip, tensile): 220 kg × 180 kg; Tear load (warp): >6524 kg.

Years ago, stretch fabrics were used exclusively in sportswear. Nowadays, they also provide added comfort in leisurewear and working apparel. The distinctive feature of clothing made from stretch fabrics is its high degree of wear comfort. Special yarns in the warp or weft, and occasionally in both directions, make the fabric elastic. It adapts to the wearer’s movements and does not go baggy where it is subjected to high tensile stress, e.g. at the elbows, knees, and seat. The degree of comfort depends on the fabric construction and on how the garment is made up.

A fabric’s elasticity is dictated by the yarn used. Originally, it was highly elastic, texturized continuous filament yarns which gave fabrics used for sportswear their stretchability. Today, it is mainly elastane threads covered with staple fibres of cotton, wool, or man-made fibres. Elastane yarns of different fineness are used, depending on the fabric and the degree of stretch required. Yarns with elastane ensure a defined degree of stretch matched to the garment and the wearer, snap-back, dimensional stability, and maintenance of stretch for the entire lifetime of the garment. Garments are considered comfortable if they have the stretch values of about 18% in the warp or weft direction in menswear and 20% in ladies’ wear.
To produce warp-elastic fabrics, corresponding adaptations have to be made during production of the warp itself. As a rule, warps consisting of elastic yarns are sectionally warped. With a high-speed warping creel, the warp can be produced with a constant and uniform tension. The warp is sized from the warping beam. In this way, the elasticity of the yarn is blocked for the weaving process, and the warp can be woven almost like a ‘normal’ warp. If elastic yarns are used in the weft, a suitable weft insertion system must be chosen, and various components of the machine designed in accordance with the yarn properties. Weft-elastic yarns need a larger reed width. To achieve adequate elasticity of the fabric, the density of warp and weft have to be defined accordingly: if the values chosen are too high, the elasticity may, in extreme cases, be blocked.

Elastic yarns can be woven on rapier and projectile weaving machines. Both machine types can be adapted to meet the special requirements for processing elastic yarns, e.g. by the use of appropriate weft feeders, weft brakes, projectiles, gripper clamps and roller surfaces. The free ends of elastic yarns always have a tendency to snap back, and therefore have to be gripped and held with high accuracy and the necessary clamping force. Besides the clamping force, special attention also has to be paid to the clamping area. Weaving systems should have an electronically well controlled warp left-off with warp tensioner and electronically controlled cloth take-up. These enable the warp tension and weft density to be maintained with great precision. It results a uniform, reproducible stretch over the entire warp length.

9.8.4 Glass fibre fabrics for PCB

Almost all devices, machines and vehicles which are powered, controlled or regulated by electricity contain printed circuit boards (PCB). These boards, which serve as carriers for the circuits, are reinforced with fabrics made of glass fibre yarns. They have to meet exceptionally high standards of quality. The fabric must be dimensionally stable, i.e. it must not exhibit any elongation in either warp or weft. The surface must be uniform and the texture should be uniform in both the warp and weft direction. The fabric surface must not exhibit any broken filaments, because broken and projecting filaments may cause the printed circuit to malfunction.

All of the glass fibre yarns used are endless multifilament yarns consisting of many individual endless threads, known as fibrils or filaments. To facilitate weaving the yarn usually has a protective twist. This makes the thread more compact and the fibrils less susceptible to damage. However, so-called ‘zero-twist’ yarns are now being used increasingly to ensure that the synthetic material penetrates the fabric layer efficiently in production of the printed circuits. The filaments in these yarns are parallel, and
therefore much more susceptible to damage and breakage. In the production of such fabrics, especially those made of untwisted glass fibre yarns, rapier weaving machines have crucial advantages as regards weft insertion. Conductive glass fibre fabrics are commonly woven on air-jet weaving machines, because weft insertion without any mechanical components minimizes wear of the fibrils. But the open thread structure of the ‘zero-twist’ yarns makes weaving problematic. Rapier weaving machines have the advantage that the open-thread yarns are securely gripped by the yarn clamps and can be inserted in the shed without any individual filaments being missed.

9.8.5 E-textiles

Electronic textiles (e-textiles) are fabrics that have electronics and interconnections woven into them, with physical flexibility and size that cannot be achieved with existing electronic manufacturing techniques. These textile structures may enable sensors and other electronic devices to be distributed over large areas in an economical manner. Some commercial electrotextile products like fabric-based heating systems, flexible keypads and keyboards, fabrics with integrated sensors, etc., have already been developed. Research in electronic textiles is currently being carried out in many diverse disciplines. It is envisioned that the preliminary research in these areas will pave the way for the development of fully integrated electronic textiles with integrated circuits, devices, and power sources, etc., built into the textile structures. Textile substrates with conducting components (fibres, yarns, etc.) are being used for the development of electrical and electronic devices and systems in applications where flexibility and conformability are of importance. Some of the research that is being carried out in this area has been applied to develop commercial products for civilian and military applications.

Applications of textile-based circuitry include a row and column fabric keyboard, an electrical dress, a musical jacket incorporating an e-broidered keypad and fabric buses, and a musical ball with e-broidered pressure sensors. One of the most significant recent developments in the application of conducting threads to electronics has been the attempt to form networks of conducting threads in a fabric by weaving. Great strides have been made in the last decade towards truly wearable unobtrusive electronic functionality into textiles. In the future electronic devices, including power supply and some level of information processing, will be built into the textile structures at various levels of fibres, yarns and fabrics. Additionally, electroactive polymer materials in some form may also be incorporated into textiles to incorporate active components.
9.8.6 Geotextiles

A geotextile is defined as any permeable textile material that is used with foundation, soil, rock, earth, etc. to increase stability and decrease wind and water erosion. A geotextile may be made of synthetic or natural fibres. In contrast, a geo-membrane is a continuous membrane-type liner or barrier. Geo-membranes must have sufficiently low permeability to control migration of fluid in a constructed project, structure or system. A geotextile is designed to be permeable to allow the flow of fluids through it and a geomembrane is designed to restrict the fluid flow.

Geotextiles have historically been made of natural plant fibre; modern geotextiles are usually made from a synthetic polymer (such as polypropylene, polyester, polyethylenes and polyamides) or a composite of natural and synthetic material. Plant fibre-based erosion control geotextiles are subject to decomposition and have a limited shelf life before their inherent durability suffers. The synthetic polymers have the advantage of not decaying under biological and chemical processes, but being a petrochemical-based product they use non-renewable resources in their construction, and cause environmental pollution in their manufacture and use, and have associated health risks.

Geotextiles can be woven, knitted or non-woven. Different fabric composition and construction are suitable for different applications. The non-woven geotextile is an arrangement of fibres either oriented or randomly patterned in a sheet, resembling felt. These geotextiles provide planar control. Woven geotextile looks like burlap. It is a fabric made of two sets of parallel strands systematically interlaced to form a thin, flat fabric. The strands are of two kinds – slit film which is flat, or monofilaments which are round. The way these two sets of yarns are interlaced determines the weave pattern that in turn determines the best application for that woven fabric. Weave patterns come in a virtually unlimited variety that do affect some properties of the fabric. Woven geotextiles are generally preferred for applications where high strength properties are needed, but where filtration requirements are less critical and planar flow is not a consideration. These fabrics reduce localized shear failure in weak subsoil conditions, improving construction over soft subsoil and providing access to remote areas through separation.

Woven monofilament geotextile fabrics provide immediate and long term solutions for most drainage and filtration applications. The combination of high strength and excellent hydraulic characteristics make monofilament the fabric of choice behind bulkheads and under riprap. Percent
open area, a property most often associated with 100% monofilament geotextiles, offers both water and particles a direct path through the geotextile. Woven geotextile fabrics are produced by weaving high tenacity yarns in an orderly pattern on large industrial looms. The high tensile strength of the order of about 200 kN/m allows the material to accommodate large stresses during installation and while in service without failure. Woven structures provide high abrasion resistance which makes the material resistant to the installation process and to sand abrasion in the surf zone. Woven geotextiles have a low capacity for elongation and consequently cannot accommodate large strains without failure. This is countered by the high tensile strength of the material, but it may lead to problems if the material is required to reshape during deployment or when in service. Woven geotextiles have a relatively low angle of friction between both the seabed and against other geotextiles, which may affect the stability of the structure as the units will be more likely to slide off one another.

9.9 Future outlook in weaving

The principle of interlacing yarns to make a woven fabric has not changed since time immemorial. However, there have been dramatic changes in the equipment used in weaving. In today’s competitive market, low-cost manufacturing of quality woven fabrics is important for survival. Despite some gains, in particular by knitted fabrics, weaving will continue to be the predominant method of fabric formation, due to the advantages of structural stability of woven fabric. Weaving machine manufacturers are finding new, faster and better ways to produce woven cloth. Over the years, electronics have increased the processing speed, flexibility and reliability of weaving machines. Multiprocessors are now used to control, monitor and communicate functions. Modern control systems are capable of generating production statistics, efficiency calculations and a variety of other data which can be retrieved from the machine via an interface. Fabric parameters, patterns, colours and control functions can be input at the communication panel on the machine. Although the speed of weaving machines has increased dramatically, the weft insertion rates and loom productivity of single-phase projectile, air-jet and flexible rapiers are expected to increase further. It is also expected that there will be major improvements in multiphase weaving as such improvements may not be possible for single-phase machines due to physical limitations. However, research programmes are under way to further improve the performance of single-phase weaving machines. For example, pneumatic beat-up is being studied as a replacement for the traditional reed in air-jet weaving. In pneumatic beat-up, compressed air is used to push the filling yarn into the cloth fell. Weaving productivity and flexibility have been substantially improved with the
introduction of the quick style change system, the off-loom take-up system, inverter drives, filling feeders, electronic let-off and take-up, automatic filling repair and new monitoring systems. Innovations to further increase productivity and flexibility are expected to continue, and may include automatic warp stop repair, automatic fabric doffing, automatic filling supply systems, etc.

With the ever-expanding fields of application of high-performance industrial fabrics, weaving machines will continue to be modified to meet the requirement of their production. Future machines would be wider and stronger to produce a wide range of industrial fabrics. One area with great potential for improvement is the manufacture of weaving machines for complex three-dimensional shapes. As industrial textiles penetrate into almost every industry in the world, 3D woven fabric structures are gaining importance. As of today, no 3D weaving machines are commercially available. However, this is expected to change as the market share of 3D fabrics increases. New fabric development will be a key factor in survival and success of the weaving industry in the years to come.

9.10 References


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10 Advances in coloration of polyester textiles

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10.1 Evolution of dyeing of polyester

Polyester fibre [poly (ethylene terephthalate) (PET)] offered considerable
difficulties in dyeing when it was first produced by the Imperial Chemical
Industries (ICI) in 1947. Since polyester has no reactive groups such as
those present in cellulose (hydroxy groups) and protein (amino groups)
fibres it had no affinity for water-soluble dyes, like direct, acid and basic
dyes that were being extensively used for coloration of textiles at that time.
Only one major class of dyestuff proved to be practical for dyeing light
shades on unmodified polyester: disperse dyes – a nonionic dye class. These
dyes were originally developed by Ellis in 1923 for the dyeing of cellulose
acetate fibres.

The first systematic study of the dyeing of polyester (polyethylene tere-
phthalate) was carried out by Waters\(^1\) in 1950. From the diffusion coeffi-
cient and saturation values of four disperse dyes dyed at 85°C and 100°C,
this investigator concluded that even though disperse dyes have an affinity
for polyester the ‘slow diffusion is the cause of the poor dyeability of poly-
ester’. In order to achieve deep shades, the dyebath temperature must be
as high as possible.

In a subsequent study, Lyle, Iannarone, and Thomas\(^2\) observed that dis-
perse (acetate) and vat dyes on polyester, when applied at 250°F (121°C),
gave an excellent penetration generally not obtained at 212°F (100°C).
Remington\(^3\) pointed out that the problem in dyeing polyester fibre results
from the compactness of the structure, which prevents dyes from diffusing
through it rapidly. Dye molecules can be induced to travel rapidly from the
dyebath to the interior of the fibre by means of carriers, use of high tem-
perature, or by forming the final dye molecules within the fibre.

 Schroeder and Remington\(^4\) were the first to propose the mechanism of
dyeing of polyester fibres with disperse dyes. According to these investi-
gators, the dye is sorbed from its solution by the polyester fibre and a constant
partition between the concentration of dye in water and the concentration
of the dye in fibre is maintained until equilibrium, thereby confirming the solid solution mechanism. These investigators further stated that since it is difficult to say whether the dye is uniformly solubilised in fibre or not (since the fibre contains crystalline and amorphous portions), the use of the term ‘solid solution’ is not appropriate.

A detailed investigation\(^5\) of the stability, dyeing, and fastness properties of a range of disperse dyes on polyester fibre showed that it can be treated for 2 h at temperatures up to 160°C in the pH range 2.8–7.0 without reducing its toughness factor. Optimum dyeing conditions were obtained between 130 and 140°C.\(^6\)

From the studies on the solubilities of the dye in water and fibre, rates of desorption from dyed fibres, heats of dyeing, diffusion coefficients, and activation energies, Patterson and Sheldon\(^7\) concluded that in the dyeing of polyester with disperse dyes the dissolution of the dye molecules in H\(_2\)O is followed by the diffusion of single dye molecules inside the fibre structure. The rate process depended on the spontaneous appearance at a suitable position in the non-crystalline regions of a hole large enough to accommodate the dye molecule.

In another similar work Glenz\(^8,9\) concluded that polyester molecules are bound together by relatively strong forces which must be loosened by the penetrating dye. Moreover, if the dye is to diffuse, it must detach itself from the site at which it is adsorbed and be anchored to a neighbouring site, with a velocity determined by the affinity of the dye for the fibre, the relative size of the intermicellar spacing and the dye molecule, the changes in energy and in geometry. To increase the diffusion rate, either the temperature must be raised or the energy of activation must be lowered, as by the addition of a carrier. According to Garrido\(^10\) the activation energy of dyeing calculated from the rates of dyeing with CI Disperse Red 15 at 70, 80, 85, 90, and 95°C using the Arrhenius equation, is about 70 kcal/mole for the dyeing of polyester fibre with disperse dyes. These pioneering studies formed the basis of the further development of dyeing of polyester.

### 10.2 Disperse dyes

The first ever ‘Disperse Dye’ SRA Orange 1 was synthesized by Holland Ellis for the British Celanese Ltd in 1923. However, the Gold Medal of the Society was given to Baddiley and Shepherdson for introducing ‘Duranol’ dyes in 1924, much before the invention of polyester fibres.\(^11\)

When polyester was introduced Duranol dyes and other such dyes were there to dye it. These dyes being simple, nitro, azo and anthraquinone dyes were found to have low sublimation fastness and those based on anthraquinone backbone were found to fade by NO\(_x\) gases found in the urban environment.
The first tasks of development of suitable disperse dyes for the dyeing of polyester were to modify the then available dyes to improve their sublimation and gas fume fading properties. The sublimation fastness was achieved by introducing additional groups in the dye molecules to increase their molecular weight and the deficiency of gas fume fading was rectified by protecting the easily reducible amino groups in the anthraquinone based disperse dyes. Subsequently, disperse dyes were developed to meet the specific requirements of polyester such as ‘alkali clearing’ dyes and the dyes having low thermo-migration and high wet fastness. The forerunners among the ‘alkali clearable’ dyes were the di-ester-side group containing Dispersol PC dyes that could be ‘reduction cleared’ without a reducing agent to remove the superficially surface adhering dye, thereby reducing the pollution load of the effluent. These dyes have also been exploited in the ‘discharge’ printing of polyester fabrics where the dye from the alkali printed portion of the printed design can be easily washed with hot aqueous alkali solution. In another approach similar results can be achieved by dyeing with thiophene-based disperse dyes, where the dye molecule is ruptured on treatment with alkali breaking the bond between the azo and the thiophene group, producing almost colourless easily washable by-products. Recently, Koh has developed alkali-clearable azohydroxypyridone disperse dyes containing a fluorosulfonyl group. The alkali-clearability is imparted by the hydrolysis of fluorosulfonyl residues and ionisation of azohydroxypyridone structure under alkaline condition.

Dyes with low thermo-migration propensity and unique wet fastness that were researched by ICI were introduced for the dyeing of polyester in 1975. These dyes such as CI Disperse Red 356 and 367 are based on benzodifuranone chromophoric backbone. Additionally, they are extremely bright with good build-up. Dyes dyeable from an alkaline medium have also been developed to cater to the needs of polyester-cellulosic blend dyeing. The traditional classification of disperse dyes on the basis of sublimation characteristics (i.e. A,B,C,D or E,SE,S) and concomitant energy levels (Low, Medium, High) has undergone considerable change. The first such change occurred when the concept of ‘Rapid Dyeing’ was introduced in the 1980s, where the ‘on-tone’ dyeing and the compatibility of dyes based on their concentration-based exhaustion behaviour was desirable. This led to the emergence of multi-dye formulated RD dyes and single-dye based compatible trichromatic dyes. In more recent times, dyes are being classified on the basis of end-use, such as dyes for automotive fabrics having very high light fastness or dyes for sportswear and heavy-duty work-wear having the highest level of wash fastness, or for that matter, luminous disperse dyes for high-visibility garments used by traffic controlling personnel. As far as home-textiles are concerned one can use compatible dyes with good levelling properties and reasonable fastness. There are various...
minor classifications that have not yet found their place in shade cards such as dyes suitable for ‘Solvent Dyeing’ and ‘SCFE dyeing’.

The disperse dyes (erstwhile Cellulose Acetate Dyes) were originally introduced as pastes that had problems of settling down and crust formation. Subsequently spray drying was introduced where a nicely ground mixture of a dispersing agent and the dye were spray dried to produce self-dispersible powders. Automation of the dyeing process again led to the production of easily ‘pumpable’ dyes in the form of liquid dyes. In the production of dyes besides their easily dispersability and pourability, the particle size is of paramount importance since the solubility of the dye is directly proportional to the particle size and hence its ultimate exhaustion of polyester. A typical dye may have particle sizes varying from 0.1 to 1 micron possibly with some nanoparticles. Studies on the crystalline structure of disperse dyes has indicated that they are polymorphic and the different crystal forms have different saturation value. However, prolonged dyeing at high temperature results in the conversion of all meta-stable forms to the stable form, hence the original difference in the crystal forms of disperse dyes does not affect the ultimate dye uptake.20,21

A new range of micro-encapsulated disperse dyes having regulated release of disperse dyes was developed in 1970 and used for creating snow-like flecked effect on fabric by printing.22 Thereafter magnetic micro-encapsulated dyes were introduced for the Thermosol dyeing of polyester in specially designed machines.23 Recently, polyurea microcapsules (PMs) having disperse dyes for Thermosol and high temperature exhaust dyeing have been produced that can dye polyester without any auxiliary chemical and the exhausted dye bath can be reused for dyeing.24

In another recent study, dispersed dyeing having temporary solubilising groups, that do not require the use of dispersing agents, have been used for the dyeing of polyester.25 In this study, precursors of vinylsulphone dyes having aminophenyl-4-(β-sulphatoethylsulphone) groups that impart water solubility to dye have been synthesised and dyed onto polyester. The β-elimination reaction of the sodium sulphate group during dyeing produces the vinylsulphone form of the dye and this in turn may be partly hydrolysed to the hydroxyethylsulphone form, both of which are sparingly soluble, during the dyeing process. The optimum application pH for the dyeing of polyester in the absence of a dispersing agent has been found to be between pH 5 and 6 at which the hydrolysis rate of the dyes is moderate. These and similar precursors of vinylsulphone dyes have been used for the single bath dyeing of polyester-cotton blends. Where the vinylsulphone form of the dye covalently reacts with cotton and is also sorbed by polyester as a disperse dye.26 This approach of synthesizing dual function disperse-reactive dyes that can dye various fibres singly or in blends is being pursued by several investigators.27
10.3 Theory of dyeing with disperse dyes

As stated above the dyeing of polyester with disperse dyes was perceived as the distribution of the dye molecules between the dye bath and polyester fibre such that at equilibrium the concentration of the dye in the polyester fibre is linearly proportional to the concentration of the dye remaining in the dye bath according to Nernst Isotherm. The ratio of the concentration of dye in fibre to the concentration of the dye remaining in the dye bath represents the Nernst distribution coefficient. This coefficient was taken as a measure of the affinity of the particular dye for a particular polyester fibre assuming it to be a homogeneous inert solvent in which dye molecules could dissolve.

However, subsequent studies on the differently pretreated polyester fibres having complex morphological structures showed that the dye distribution in the fibre was different for fibres having different morphology even though it had the same basic chemical structure. These observations led to the closer examination of the state of the dye in the dispersion, particle size and crystal structures, morphology of the fibre, its previous history and changes during dyeing, and diffusion in fibre.

While deliberating on the state of the dye in the dye bath McDowell postulated that the dyeing of polyester fibres with disperse dyes takes place from dilute solution of dye in the presence of a suspension of dye particles. As the dye molecules are sorbed by the fibre from solution, more dye passes into solution from suspension in order to maintain the concentration of the solution, which is in equilibrium with a definite concentration of dye on the surface of the fibre. As dyeing proceeds and these molecules are taken up from solution by the fibre, the dye remaining in suspension in the form of much larger particles dissolves slowly and is taken up by the fibre until equilibrium is reached as shown in Fig. 10.1. When equilibrium is reached, the following subsidiary equilibria are established.

I. Dyeing dispersion in bath ⇔ Dye dissolved in the bath
II. Dye dissolved in the bath ⇔ Dye dissolved in the fibre
III. Dye dissolved in the fibre ⇔ Dye diffused in the fibre

Theoretical studies of the equilibrium adsorption of dyes showed that two types of isotherms are obtained, viz. (1) Isotherm indicating Langmuir type adsorption of dyes, and (2) Isotherm indicating constant partition ratio up to the saturation point.

From the studies of Patterson and Sheldon it was concluded that in the dyeing of polyester with disperse dyes the diffusion of single dye molecules inside the fibre structure takes place. Moreover, the rate process depended on the spontaneous appearance at a suitable position in the non-crystalline regions of a hole large enough to accommodate the dye molecule. These
findings have led to the postulation of Free Volume Theory of diffusion of dyes into thermoplastic fibres that is applicable to the disperse dye polyester fibre system.\textsuperscript{32} According to this theory the dyestuff is attracted from solution to the fibre surface and remains there until sufficient thermal energy has been supplied to the system to allow the dyestuff to diffuse into the fibre through the transient holes, the formation of which is a result of the onset of segmental motion of the fibre. Thus at and above the glass transition temperature (Tg) of the fibre the dyestuff diffuses into the fibre by ‘jumping’ from one site to another (Fig. 10.2).
Below the glass transition temperature, the polymer molecules are relatively motionless and very little, if any, diffusion of dyestuff into the fibre takes place. Once the T_g is reached, sufficient energy exists in the fibre causing some segments in the polymer to move. These segments vibrate, but no net translational motion of the centre of the molecule occurs. When a polymer chain comes in contact with a chain having no dye molecule, the dyestuff can interchange between these two chains and thus diffuse through the polymer.

In the free volume theory, the activation energy of dyeing is defined as the energy required to form a free volume around the dye which can accommodate the dyestuff and permit a location change of the incorporated dye molecule. The activation energy is not constant but is a variable with a maximum value at the T_g, decreasing with increasing temperature. The purpose of the solvent, such as water, according to the free volume theory, is to transport the dye from solution to the fibre surface and then to act as plasticiser. The mode of action of the plasticiser in dyeing is to lower the energy of activation. The plasticiser is sorbed into the fibre and weakens the intermolecular attractive forces. Due to weakening of these forces, the plasticiser energy needed to permit rotational and vibrational motion of the chain segments is reduced, thus lowering the T_g of the fibre. The concentration of the plasticiser in the fibre is more important than the chemical structure of the plasticiser.\(^{33}\)

In this theory of diffusion of disperse dyes into polyester fibre considerable importance has been given to the T_g of the fibre and it is assumed that irrespective of the dye used the dye-diffusion starts at T_g. The ‘energy effects’, that is, the intermolecular forces between the dye and the fibre and geometric dimensions, that is, size and shape of the diffusing molecule, have not been considered.

Gulrajani and Saxena\(^{34}\) have indicated that practically no diffusion of the dye starts at the T_g of the fibre. The diffusion of the dye starts at a temperature T_D which is at a higher temperature than T_g, so that,

\[
T_D = T_g + \Delta T_{\text{Dye}}
\]

\(\Delta T_{\text{Dye}}\), the difference between \(\Delta T_D\) and T_g, has been attributed to the ‘energy effects’ as well as ‘geometric properties’ of the dyes. The value of \(\Delta T_{\text{Dye}}\) is different for different dyes. The observation that the plasticisers such as water and other organic chemicals reduce the T_g of the fibre was further investigated by Mitsuishi and coworkers.\(^{35}\) These investigators observed that presence of the dye in the fibre lowers the T_g in a similar manner as the plasticisers and is proportional to the concentration of the dye in the fibre. The lowering of T_g by the dye is influenced not only by
the molecular structure of the dyes but also the dye-polymer interactions. This observation shows that the Tg of the fibre goes on changing during dyeing.

Most of the investigators support the hypothesis that a sorption of disperse dyes on polyester follows a linear relationship between dye in the bath and dye on the fibre and dye diffusion takes place by the segmental motion of the polymer chains as postulated in the Free Volume theory. However some workers obtained Langmuir types of sorption isotherms on equilibrium dyeing of disperse dyes on polyester fibres. Such curves are associated with processes in which the dye is attached to the sites on the surfaces of crystallites or to the sites on the molecular chains in the amorphous region.

It is reported that the sorption isotherms depend on the presence of dispersing agents, the dyeing temperature, and the dye structure and fibre types. Generally, sorption isotherms of disperse dyes on polyester fibres have been reported as Nernst-type linear distribution isotherms in the presence of a constant concentration of dispersing agent. However, in the absence of a dispersing agent, the sorption isotherms of disperse dyes on polyester fibres have more complex shapes. It has been reported that a dual-mode sorption model that is composed of both Nernst and Langmuir sorption models, or a more complex sorption model which has one Nernst type model and several Langmuir sorption models, may be more appropriate.

Nakamura et al. while investigating the sorption isotherms and dyeing rates of purified disperse dyes on polyester microdenier fibres from aqueous medium obtained isotherms that are curved and can be described in terms of the dual-mode sorption model: Nernst-type partitioning and Langmuir sorption isotherms. However, this dual-model also depends on the dye structure and on the dyeing temperature. Therefore, the selection of the sorption isotherm model for polyester disperse dyeing systems depends on dyeing and material conditions.

While investigating the diffusion of disperse dyes into microfibres and conventional polyester fibres, Park et al. observed that the simple Langmuir sorption isotherm fits well to the experimental sorption isotherms obtained at high dyeing temperatures. The correlation is, however, better for microfibres than for conventional fibres. These findings of Park et al. have been very recently confirmed by Dhouib et al. According to these researchers the most important parameters for dye saturation on the fibre are temperature, dye size and volume of accessible domains able to receive dye molecules. This volume is very dependent on fibre crystallinity. Both fibre surface area and dyeing temperature have a second-order effect. The temperature has more influence on fibres having poor accessibility to the dye.
10.4 Effect of fibre structure on dyeing

10.4.1 Heat setting

In the production of polyester fibre, variations in crystallinity, orientation of the crystalline, paracrystalline and amorphous regions of the fibre may occur during the post-spinning drawing stage and the heat-setting processes. These changes in the physical structure of the fibre affect the dye-ability of the fibre. Several attempts have been made to establish qualitative and quantitative correlation between the dye diffusion and morphology of polyester fibre. Marvin\textsuperscript{42} was the first to show that over a range of setting temperatures the uptake of disperse dyes by heat-set polyester initially decreased, as the temperature of pre-heating was raised giving minima in the range 140°C to 180°C. However, at higher temperatures the dye uptake increased with temperature and was greater than that of the untreated control sample. These observations of Marvin have been confirmed by many other workers.

The anomalous dye uptake of heat-set polyester has been explained by Dumbleton \textit{et al.}\textsuperscript{43} on the theory that the diffusion of dye depends on its segmental mobility in the amorphous regions and also on the size and size distribution of the crystallites. Moreover, it is also influenced by the number of crystalline sites and the orientation of the amorphous phase within its fine structure. Some workers have observed that the porosity of the fibre, that is, the number, size and size distribution of pores within the matrix, can also affect dye uptake.\textsuperscript{44–47} In a comprehensive study\textsuperscript{48} of the effect of polyester fibres heat-set in slack and taut condition on the diffusion behaviour of disperse dyes, it has been proposed that dye diffusion in heat-set polyester is controlled by two factors: (1) the volume of the accessible region, represented by the amorphous volume per crystal; and (2) the tortuosity of the path of diffusion of dye into the fibre that is dependent on the orientation of the amorphous region and the type of coupling between the amorphous and the crystalline region. Donze \textit{et al.}\textsuperscript{49} have suggested that adequate shrinkage must be allowed during heat-setting of polyester fabrics to smooth out the structural differences caused by variations of the tension applied to the yarns during mechanical textile processes.

On dyeing of polyester heat-set at temperature between 160°C to 220°C with disperse dyes at 130°C Gacèn \textit{et al.}\textsuperscript{50} observed that the fibre structure becomes more compact, thereby showing that supplementary setting of polyester takes place in the dye bath at 130°C. This effect is more pronounced in fabrics heat-set at relatively low temperatures. These findings contradict the earlier finding of Lipp-Symonowicz\textsuperscript{51} that treatment of polyester fibres in water at 130°C decreases the molecular orientation of polymer
chains in the surface layers of the fibre. However, both these studies show that some further setting of the polyester takes place during dyeing.

10.4.2 Texturing

Texturing of polyester filament yarns not only disturbs the orientation of the polymer chains and the crystallinity of the yarn but it also modifies its cross-section resulting in uneven dye sorption by the yarn thereby producing barré dyeing. Clements\textsuperscript{52} devised a method of evaluating the dye uniformity of false-twist pin-textured polyester yarns produced from fully drawn feed yarns. According to this investigator, the textured yarn dyeability is not very sensitive to the texturing parameters, and only a serious lack of drawback of proper process control can account for very high dye variance of some commercial yarns.

From a study of the dyeability of poly (ethylene terephthalate) yarn textured at different temperatures and different contact times Gupta, Kumar and Gulrajani\textsuperscript{53} have concluded that the degree of crystallinity and the size of crystallites are the controlling factors of dye uptake. Moreover, the temperature of texturing determines the crystallinity and the crystal size distribution in the textured samples. Their findings have been supplemented by the observations of Goldin.\textsuperscript{54} According to this investigator the nonuniformity of dyeing textured yarn is related to the shape of the primary heater response curve in texturing. According to Miller, Southern and Ballman\textsuperscript{55} the variable dyeability of textured yarn that is dependent on yarn crystallite size and crystalline content can be attributed to dye path tortuosity of disperse dyes in textured polyester fibres.

In a comprehensive study of the response of textured yarn properties to process variables in relation to barré, McGregor and coworkers\textsuperscript{56} have concluded that the rate of dyeing is not very sensitive to texturing variables, and the control of the heater temperatures should be within ±2°F (1°C) to get barré free dyeings and the feed yarn should be uniform. In a subsequent study,\textsuperscript{57} they have reported that differences in feed-yarn orientation propagate though texturing and modify almost all the textured yarn properties. Such effects can be comparable in magnitude with those of large changes in texturing variables, and they influence some of the crimp, shrinkage, and tensile properties of the yarns even more than they modify the dyeing responses. However, the information on feed yarn variation is lost on high-temperature dyeing.

Navrati\textsuperscript{58} has proposed a mathematical model to predict disperse dye barréness attributable to differences in draw ratio and texturing temperature of polyester fibres. It is claimed that the use of dyeing simulation will be useful to optimize dye formulations and dye usage. In an another study\textsuperscript{59} it is proposed that determination of crimp contraction force
variations is an effective way of detecting potential nonuniform dyeing behaviour of textured yarns. When the coefficient of variations in crimp contraction force and percentage tension instability were >5%, barré appeared more frequently in textured yarn fabrics so that a strict tension control during draw texturing is necessary to avoid barré problems.\textsuperscript{60} Lawson-Hemphill has developed Textured Yarn Tester that employs dynamic testing of crimp and can be used for determination of dyeability properties of polyester textured yarns.\textsuperscript{61} ASTM standard D 6774-02 titled ‘Standard Test Method for Textured Yarns Using a Dynamic Textured Yarn Tester’ specifies the use of this tester.

### 10.5 Dyeing procedures

#### 10.5.1 Exhaust dyeing

A conventional dyeing cycle has three main phases, namely, the sorption phase, the diffusion and levelling phase and the after-treatment phase. The levelling phase that is about 45 min to 60 min takes care of the uneven sorption of the dye during the sorption phase. This phase is a drain on energy and productivity. For the dyeing of polyester it was soon realized that unlike natural fibres the fibre plays an active role in dyeing. Hence the rate of sorption of dye suddenly jumps above the Tg of the fibre when the segmental motion sets in, giving rise to rapid diffusion of the dye into the fibre. This necessitated the need to work out a differential heating up procedure for the dyeing of polyester. In doing so it was observed that if the sorption of the dye onto fibre is regulated during the first phase of dyeing then the need for a levelling phase is reduced or even eliminated in some cases. This gave birth to the Rapid Dyeing of polyester.\textsuperscript{62–67}

The rapid dyeing of polyester, based on the differential heating rate from 60°C to 130°C, resulted in considerably shortening the dyeing time with the result that new rapid dyeing dyes and machines had to be developed. Every dye manufacturer came out with their own heating up procedure and set of dyes that was suitable for that procedure. Over time, the situation has improved thanks to the introduction of programmable microprocessor, attached to the dyeing machines that could carry out different kinds of dyeing protocols from its stored memory.

A further development in the dyeing of polyester is that many companies have produced dedicated software suitable for their own range of dyes such as the Optidye P of DyeStar for Dianex CC dyes. These software claim to optimize various parameters of dyeing and reduce the auxiliary consumption. BASF has developed its own integrated polyester dyeing system ‘dyexact XP’, which enables precise control of the dyeing process.\textsuperscript{68} The essential core product of dyexact XP range, Basojet XP, has been designed
to cater for all 100% polyester exhaust dyeing, of which 70–80% can be processed with this product alone, in combination with normal pH control. According to BASF, Basojet XP, the basis of dyexact XP range, is an effective levelling and disposing agent with excellent anti-oligomer property suitable for high temperature dyeing of polyester with disperse dyes. It offers a large number economic and ecological advantages such as process control and high levels of process reliability.

Besides these some dye-bath exhaustion monitoring software based on Flow Injection Analysis (FIA) methods to monitor the dyeing of polyester, cellulose acetate and cotton have been developed. These methods avoid some of the problems of direct spectrophotometric dye-bath measurement of dye concentrations. Using FIA methods successfully monitored the disperse dyeing of polyester yarns and the individual exhaustion characteristics of three mixed dyes can be carried out.

10.5.2 Continuous dyeing

A process for continuous dyeing of polyester by padding the fabric with disperse dyes and curing the padded, dried fabric at 180–220°C for 45–180 s was developed and patented by du Pont in 1953. This process which is called ‘Thermosol Dyeing’ was licensed for use by industry for the dyeing of polyester-cellulosic blended fabrics in the late 1950s. Since then the process is being mainly used for dyeing of polyester blends on continuous dyeing ranges (CDRs).

Numerous studies have been carried out on the optimization of the degree of fixation as a function of the concentration of the padding bath and of the temperature and time of heat fixation in the Thermosol dyeing of polyester fibres. It is reported that the rate of diffusion of dye into polyester is doubled for a 10°C temperature increase.

The use of dyes having good sublimation fastness is recommended. However, it has been observed that the higher the sublimation fastness of the dye, the higher the temperature required for its levelling and the higher the operating temperature required in the dyeing; hence, the choice of dyes employed in mixtures is important for attaining maximum dye buildup. From the studies on the Foron S and SE dyes of Sandoz (now Clariant) Somm and Gerber have surmised that the diffusion coefficients of dyes with low sublimation fastness are greater than those with high sublimation fastness but during dyeing the former are absorbed through vapour phase migration while the latter are absorbed by direct contact.

Attempts have also been made to dye both cotton and polyester with vat dyes. Vat dyes generally give different shades on polyester; however, some of the dyes were found to give reasonable good shades. In a recent study it has been reported that colour depth of vat dyes increased by adding
urea in the padding solution.\textsuperscript{74} The use of auxiliary chemicals in the padding bath to increase the diffusion and uptake of disperse dyes has been extensively investigated. Since the Thermosol dyeing of polyester proceeds by first dissolving the dye on the surface layer at low temperature followed by its diffusion into fibre at high temperature, those compounds that have high solubilising capacity and melt during curing can increase the dye uptake.\textsuperscript{75} Many nonionic surfactants have been recommended as accelerants for intensifying the colour and yield of disperse dyes applied by Thermosoling process. Their mechanism of action has been recently reviewed by Manian and Etters.\textsuperscript{76} According to these authors nonionic surfactants may act as fixation accelerants by accelerating the rate of dye dissolution in the auxiliary melt. At high surfactant concentrations, dye retention in surfactants may adversely affect dye fixation.

Decheva \textit{et al.}\textsuperscript{77–79} on the basis of their intensive study have recommended the use of caprolactam as an additive in the padding bath for its intensifying effect. According to these investigators the solubilising and dispersing effect of caprolactam, its chemical inertness toward disperse dyes, its non-detrimental effect on colour fastness and fibre properties, its ability to decrease the temperature and time of dyeing, and its positive effects on fixation and other properties of a wide variety of disperse dyes make it an ideal intensifier for use in Thermosol dyeing of polyester fibres. The use of caprolactam results in lowering the Thermosol temperature by 10˚C and Thermosoling duration by 15 s. In a later study Dusheva and Aleksandrova\textsuperscript{80} have stated that the presence of caprolactum enhances the effect of temperature on the supramolecular structure of polyester by increasing segment mobility of the macromolecular chain in the non-crystalline regions, and facilitates the relaxation and crystallization processes so that a supramolecular structure of higher micro-heterogeneity and, correspondingly, better accessibility of the non-crystalline area of the fibre is created. These changes facilitate the formation of new crystalline aggregates under the effect of disperse dyes. Caprolactam also decreases the oligomer content on the fibre surface and creates an energetically more uniform surface which promotes sorption. The rate of dyeing increases 1.5-fold.

While investigating the effect of orientation of the fibre on the rate of dyeing it was observed that rate of Thermosol dyeing was much greater in partially elongated than in fully elongated fibres.\textsuperscript{81} On the other hand it was observed that the orientation of the fibre decreases with increasing Thermosoling temperature.\textsuperscript{82} Moreover, the fixation of disperse dyes on polyester fibres in Thermosol dyeing decreases with increasing crystallinity of the fibres.\textsuperscript{83} However, on Thermosol dyeing, three types of crystalline aggregates are formed, that is, aggregates containing only polymer molecules, those containing predominately polymer molecules but
incorporating some dye molecules, and those the recrystallization of which has been influenced by the dye molecules incorporated in the crystal lattice.\textsuperscript{84}

\textbf{10.6 New methods of dyeing}

\textbf{10.6.1 Non-aqueous dyeing}

\textit{Organic solvent dyeing}

Dyeing of polyester from aqueous medium generates substantial effluent that needs to be disposed of after proper remediation treatment. Attempts were made during the 1970s to replace water with organic solvent for the exhaust as well as continuous dyeing of polyester. Of the various solvents investigated perchloroethylene and trichloroethylene were found to be promising. The economics of the process depended on the recovery of the solvent (∼95\%) from the exhausted dye bath after dyeing.

Exhaust dyeing of polyester from perchloroethylene with disperse dyes results in higher dyeing speeds due to faster diffusion, better levelling and migration and better oligomer removal. However, the exhaustion of the dye is low and addition of auxiliary solvents such as DMF, methanol and benzyl alcohol in the presence of 1\% water in the dye bath increased it to almost 100\%.\textsuperscript{85–87} These auxiliary solvents and water have been reported to lower the \textit{Tg} of the fibre and also act as solvents for the dye.\textsuperscript{88}

In another study it is postulated that in solvent dyeing of polyesters with disperse dyes, diffusion and partition coefficients are controlled through choice of the solubility parameters of the solvent. If a solvent is selected that has a solubility parameter close to that of polyester the diffusion coefficient will be high and the partition coefficient will be low. If a solvent is selected with a very different solubility parameter from that of the polymer, the diffusion coefficient will be low while the partition coefficient will be high.\textsuperscript{89}

In a recent study by Kim, Son and Lin,\textsuperscript{90} the adsorption and solubility properties of disperse dyes on polyester from 29 solvent were examined. They observed that the dye adsorptions in alkane media were much higher than those in other non-aqueous systems. From the results of the relationship between the adsorption and the solubility of disperse dyes, it was concluded that the dye adsorptions onto polyester were linearly and inversely proportional to the dye solubilities in non-aqueous media. In a subsequent study with ten different dyes dyed from pentane Kim and Son\textsuperscript{91} confirmed their previous observation about the relation between dye sorption and its solubility. While investigating the interaction of polyester with various solvents it was observed that some of the solvents modified the
physical structure of the fibre resulting in higher dye uptake.\textsuperscript{92} This observation led to a series of studies on the dyeability of solvent pretreated polyester at or below 100°C.\textsuperscript{93–97} Weigmann \textit{et al.}\textsuperscript{93} have thoroughly investigated the effect of DMF on polyester and proposed that the treatment of polyester with DMF at high temperature results in the swelling of the fibre that leads to the formation of crystallites within the swollen structure, which, depending on the size and stability of the crystallites, can be stabilized to form voids upon removal of the interacting medium. A rigid pore mechanism of dye diffusion operates in this structure as opposed to the free volume mechanism of diffusion of dyes in thermally treated polyester yarns. In a later study\textsuperscript{94} these investigators have claimed that the solvent treatments of polyester yarn, which increase dyeability, do not change the dye-diffusion mechanism and the significant increase in dye-diffusion coefficient resulting from the solvent treatment is attributable to increased segmental mobility in noncrystalline domains of the treated fibre. The increased amount of dye is believed to be held in voids in the fibre structure formed during solvent treatment. Pretreatment in DMF also resulted in a lowering of tensile strength and initial modulus, whereas subsequent dyeing caused further reduction in extensibility.\textsuperscript{95}

Moore and Weigmann\textsuperscript{96} while studying the effect of pretreatment of polyester fibres with CH\textsubscript{2}Cl\textsubscript{2} found that this pretreatment led to greater dye penetration, due either to residual CH\textsubscript{2}Cl\textsubscript{2} in the yarn or to solvent-induced structural changes occurring in the yarn during pretreatment. The literature on solvent-induced modifications of polyester structure, properties and dyeability has been reviewed;\textsuperscript{97} however, no commercially successful process had been adopted at the time of writing.

10.6.2 Dyeing in supercritical carbon dioxide

A process of dyeing from supercritical carbon dioxide (scCO\textsubscript{2}) was patented by Schollmeyer \textit{et al.}\textsuperscript{98} in 1990 with a claim that when a polyester knitted fabric is dyed with the disperse dyes from supercritical CO\textsubscript{2} at 130°C and 250 bar pressure the dyeings were more brilliant, had a 20–40\% increase in depth of dyeing, and were comparable in fastness properties to those produced by the conventional Thermosol process. This process offers a number of ecological and economic advantages, such as, no treatment of wastewater, no drying, improved dye utilization, no addition of chemicals, no reductive after-treatments, shorter dyeing times, and no restrictions on location because of availability of water.\textsuperscript{97}

Various studies have indicated that scCO\textsubscript{2} does not damage or modify the fibre during dyeing provided it is properly heat-set preferably at a
temperature that is 30˚C higher than the dyeing temperature. The Tg of the fibre is not significantly altered in scCO₂ hence the dyeing should be carried out at 90˚C to 140˚C as in the case of conventional HT-HP dyeing. Higher temperature should be used for darker shades.\textsuperscript{100}

During the scCO₂ dyeing of polyester disperse dyes are dissolved into the supercritical phase, transported to the fibre and adsorbed onto the surface. Finally, the dye molecules diffuse into the CO₂-swollen polymer matrix, where they are held mainly by dispersion forces. On reducing the pressure after dyeing, the CO₂ molecules escape from the fibres while the dye molecules are retained. This dyeing mechanism was suggested by Saus \textit{et al.}\textsuperscript{101} and subsequently confirmed by Tabata \textit{et al.}\textsuperscript{102} ScCO₂ has a density and dissolvability similar to liquids, but viscosity and diffusion properties similar to gases.

Suitability of disperse dyes, that have been developed for dyeing from an aqueous medium, for dyeing from scCO₂ has been extensively investigated.\textsuperscript{100,103} Generally solubility is low for all dyes and is between $10^{-4}$ and $10^{-7}$ mol dye/mol CO₂ under the dyeing condition of 120–140˚C and 300 bar pressure which is similar to that in water. Moreover, the dye uptake in CO₂ is nearly comparable to conventional aqueous dyeing but not to organic solvent dyeing.\textsuperscript{102} It has been observed that changes in the temperature and pressure result in significant colour differences in terms of colour yield and hue. During dyeing, the dyes seem to compete for accessible regions of the fibre where the dyes with the lowest molecular weight have the highest mobility resulting in higher dye uptake compared with the higher molecular weight dyes. This causes differences in colour with slight variation in temperature and pressure of dyeing, especially when mixtures of dyes are used.\textsuperscript{104} Keeping the non-linear exhaustion behaviour of the dyes in scCO₂, a new colour matching system has been developed.\textsuperscript{100}

First dyeing machine with a capacity to dye 4 bobbins of 2 kg yarn each was constructed by Jasper GmbH & Co in 1991 and installed by Amann & Söhne GmbH & Co. However, there was need to improve the technology. Later on UHDE Hochdrucktechnik GmbH of Germany made a pilot plant having improved technology that also has some shortcomings.\textsuperscript{100} A commercial-scale 1000-litre supercritical dyeing machine has been designed, for treating 300 kg polyester while recycling all dye and 96% of the CO₂ by van der Karaan.\textsuperscript{105} An economic analysis showed that, although the purchase cost for a supercritical machine is higher (k€ 500) than for an aqueous machine (k€ 100), the operating cost is lower (0.35 instead of € 0.99 per kg polyester). This is due to the higher rate of dyeing and by the simpler dye formulations that can be used in scCO₂. The overall result is a 50% lower process cost for the supercritical process.
10.6.3 Dyeing from alkaline medium

Dyeing of polyester from an alkaline medium offers an opportunity to dye polyester cellulosic blends by a single bath single stage process. Most of the disperse dyes are adversely affected when heated in an alkaline bath; however, some select dyes have been found to be stable in an alkaline dye bath. It has been observed that the majority of disperse dyes are sensitive to the water hardness and the residual impurities present in the fibre when dyed in alkaline medium. Moreover the pH of the dye bath shifts towards neutral during dyeing.\(^\text{106}\) In order to overcome these problems of dyeing in an alkaline medium a special auxiliary chemical such as Diaserver AD-95 (DyStar) that has many functions (e.g. in stabilizing dyestuffs, sequestering agents, buffering, and dissolving oligomers) must be used. Optimal conditions for attaining higher \(K/S\) values for alkali stable Dianix AD dyes of DyStar have been worked out to be 130˚C for 45 min in the presence of 2% Diaserver AD-95.\(^\text{107}\)

The alkaline dyed polyester showed increased values of specific breaking stress and elongation, but lower abrasion resistance than the acid dyed yarns. The results of the extraction of oligomers showed that the virgin and the acid dyed polyester samples possessed higher oligomer content, whereas the oligomer content in the alkaline dyed polyester was lower. The measured colour intensity of the samples dyed with the Dianix AD dyes in an alkaline medium was slightly lower than that of the acid dyed samples.\(^\text{108}\)

10.7 Dyeing of chemically modified polyester fibres

10.7.1 Basic-dyeable polyester

Chemical modification of PET polyester permits a substantial change in its dyeability without any significant change in their physicomechanical properties. Of the various methods of modification, incorporation of 2–3 mol % of sodium salt of 5-sulphetoisophthalic acid into the basic PET polymer to impart basic dyeability has been commercially successful and various anionically modified fibres have been introduced in the market. The possibility of dyeing with cationic dyes entails the following advantages compared with normal disperse dyeing: (a) dyestuff costs are lower; (b) shades are more brilliant; (c) reduction clearing step is eliminated; and (d) sublimation fastness of the dyeings is high. There are some problems that need to be looked into, such as: (a) light fastness of the cationic dyes; (b) levelling and reproducibility of the dyeings due to very high affinity of the dyes for these fibres; and (c) hydrolysis of the fibre during dyeing.\(^\text{109}\)

Since these fibres are prone to hydrolysis during dyeing they are dyed at 120˚C in the presence of 2–6 g/L of sodium sulphate to protect the fibre.
It is postulated that the addition of salt prevents the exchange of sodium ions of the fibre with the hydrogen ions that is the critical step in the hydrolysis of the fibre.\textsuperscript{110}

The cationic dyes form electrovalent bonds with the sulphonic acid and carboxyl groups of the fibre and the amount of dye sorbed by the fibre is dependent on the number and accessibility of these groups. The accessibility of the anionic groups is a function of both the morphology and temperature of dying. X-ray diffraction data showed that the crystalline regions of these fibres are formed of segments of unaltered poly (ethylene terephthalate); the modified polyester segments remained in the noncrystalline regions. The degree of crystallinity of the modified fibres of similar thermal history is 10\% lower than that of the unmodified fibres and the density and orientation of noncrystalline regions in the modified fibres is lower than in the nonmodified fibres.\textsuperscript{111} Ingamells, Lilou and Peters\textsuperscript{112} have reported that the accessibility of the sulfonate groups in the fibres is greatly improved by plasticisation and is independent of the size of the exchanging ion. Since the fibre structure limits the accessibility of anionic centres so the sorption of the dye is not in stoichiometric proportion to the dye sites.

Pal, Gandhi and Kothari\textsuperscript{113} produced cationic dyeable polyester (CDPET) having different amounts of dimethyl ester of 5-sulfoisophthalic acid (DMS) salt through melt blending of normal polyester and cationic dyeable polyester chips. The presence of DMS salt disturbs the structure of CDPET fibre. The dyeability of both disperse dyes and cationic dyes on CDPET improves with increase in DMS salt content. The proper amount of DMS salt in CDPET is important to get a fibre with good dyeability as well as satisfactory mechanical properties. The mechanical properties of textured CDPET with more than 1.5 mol \% DMS salt decrease rapidly, indicating the limit of increase in DMS salt in CDPET fibre.

10.7.2 Easy-dyeable polyester

A large number of processes for the modification of polyester by incorporating a comonomer or copolymer into the PET so as to make it dyeable at or below 100\°C have been patented over 40 years.\textsuperscript{114–117} Such fibres are collectively termed as easy-dyeable polyesters. Such polyester fibres have low crystallinity and low Tg. Due to their having low Tg the segmental motion of polymer chains in the dye-accessible regions sets in at lower temperature thereby allowing the dye to diffuse at temperature below boiling. Dyeing properties of two recently introduced polyester fibres, that is, an easy-dyeable polyester (EDP) fibre and polytrimethylene terephthalate (PTT) fibre, has been investigated by Kim, Son and Lin.\textsuperscript{118} EDP is manufactured by polymerizing terephthalic acid with a mixture of polyethylene glycol and normal ethylene glycol. In the case of polyethylene
glycol, the degree of polymerization is 500–1000. This approach for EDP fiber is to relax the compact structure of the fiber molecules, to improve its hydrophilicity and finally to reduce its dyeing temperature. In addition, PTT is produced by polymerizing terephthalic acid with 1,3-propandiol instead of ethylene glycol to increase mobility and flexibility of the polymer molecular chains and to make it easy for disperse dye to penetrate into the fibre substrates. They found that the standard affinities of EDP and PTT were a little higher than those of PET and the activation energy of EDP and PTT was lower than that of PET thereby indicating that these fibres can be dyed at lower temperatures than PET.

One approach to produce deep-dyeable polyester involves the production of polyester from the PET/SiO₂ nanocomposites. The TEM photographs show that the SiO₂ nanoparticles get dispersed in the PET matrix at a size level of 10–20 nm. The DSC results indicated that the SiO₂ nanoparticles acted as nucleating agent, promoting the crystallization of the PET matrix from melt but inhibiting the crystallization from the glassy state, owing to the ‘crosslink’ interaction between the PET and SiO₂ nanoparticles. Dyeing of such fibres gives deeper dyeings.

### 10.7.3 Surface modification of fibres

For the past 25 years, studies are being pursued on the surface modification of polyester to enhance their dyeability and depth of colour by exposing them to plasma, UV laser and vacuum UV radiations in different gaseous atmospheres. Four approaches are being pursued to get deeper colours on polyester, namely: (a) optical effect due to nano-roughening of the fibre surface that modifies the reflectance of light from the surface; (b) changes in the morphology of the exposed surface; (c) creation of additional functional groups; and (d) anti-reflective film coating of the surface.

Raffaele-Addamo et al. observed an increase in colour depth upon dyeing plasma treated polyester fabrics. This has been attributed to optical effects caused by the plasma-induced increase of surface roughness that decreases the fraction of light reflected from treated surfaces with respect to smoother surfaces. Other effects such as increased surface area and modifications of the partition equilibrium of the dye between the dyeing bath and the macromolecular surface may also play a role.

It is reported that exposure of polyester fabrics to UV laser causes regular ripples perpendicular to the axis of the fibre. The amorphous regions of the surface as well as the number of carboxyl groups increase after UV laser treatment. A greater depth of shade was achieved on treated fabrics compared with untreated fabrics dyed with the same amount of disperse dye. This is due to the scattering of light caused by ripples on the
fibre surface, and greater dye uptake by the amorphous regions on the surface of laser irradiated PET fabrics.

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11.1 Background

Polymeric materials are a large and growing fraction of the fire load in homes, in commercial environments and in transportation, and are inherently flammable. In fact, in the majority of real fires, some type of plastic material is implicated, either as the primary fuel source or as one of the main contributors to the spread and severity of the fire. In addition, fires involving these materials are considerably more smoky, which is a real concern as it is now well established that smoke inhalation accounts for over 80% of all fire fatalities. Flammability, which generally refers to the propensity of a substance to ignite easily and burn rapidly with a flame, is only one indicator of fire hazard. The heat release rate (the total as well as the peak heat release rate) is the critical variable that governs the true severity and intensity of a real fire, thus dictating its destructive potential in terms of human loss and property damage.

Flame or ignition resistance is only a low level indicator of the fire safety of a material, which can be achieved with commodity plastics by adding flame-retardant chemicals. Consequently, the economic incentive to add flame retardants to commodity polymers to pass certain prescriptive flammability tests has focused polymer flammability research over the past few decades on the mechanisms and efficacy of flame-retardant additives, rather than on polymer flammability as an intrinsic material property. This trend in research, combined with the fact that flaming combustion of solids is a highly coupled, multi-phased process, and that the results of different fire tests depend on the apparatus, test conditions and sample geometry, has limited our understanding of polymer flammability.

A number of strategies, such as fuel control, fire detection, and smoke management and suppression, are continuously implemented to reduce the burden of fire. Of these, fuel control has the greatest potential to minimize fire losses. This can be achieved either by altering materials via fire retardants or by designing materials explicitly to minimize their adverse reac-
tion to fire. Traditionally, the rate of heat release has been considered the key parameter in fire safety engineering; nevertheless, ignition is also seen as the parameter that affects fire initiation, flame spread and fire growth. Remote ignition is, for example, the fastest mode of fire spread and can only be achieved by radiation heat transfer from the fire. Furthermore, flame spread can be in general described as a sequence of piloted ignitions.

11.2 Introduction

Organic polymers degrade to give volatile combustible materials when they are heated above certain critical temperatures, which in turn depend on their chemical structures. If the gaseous mixture resulting from the mixing of degradation volatiles with air is within the flammability limits, and the temperature is above the ignition temperature, then combustion begins. The combustion of a polymeric material is a highly complex process involving a series of interrelated and/or independent stages occurring in the condensed phase and the gaseous phase, and at the interfaces between the two phases (Cullis and Hirschler, 1981).

Successful strategies to reduce flammability of a polymeric material involve interrupting the complex stages of the combustion process at one or more points so as to reduce the rate and/or change the mechanism of combustion at that point. From a practical point of view, this is achieved either by the mechanical blending of a suitable flame-retardant compound with the polymer substrate (i.e. by introducing an additive) or by chemical incorporation of the retardant into the polymer molecule by simple co-polymerization or by the chemical modification of the preformed polymer (i.e. using a reactive component).

Both additives and reactives can interrupt the burning cycle of a polymer in several ways: by altering the thermal decomposition mechanism of a polymer; by quenching the flame; or by reducing the heat transferred from the flame to the decomposing polymer. The flame retardant can also cause a layer of carbon to appear on the surface of the polymer undergoing combustion. This may occur through a dehydrating action of the flame retardant, generating unsaturation in the polymer. The unsaturated structures form a carbonaceous layer by cyclization and cross-linking. The higher the amount of residual char after combustion, the lower the amount of combustible material available to perpetuate the flame and therefore the greater is the degree of flame retardance of the material. Consequently, one of the ways to achieve high degrees of flame retardancy or non-combustibility of polymeric materials is to increase the amount of char production on combustion.
Successful additives include: (a) polyhalogenated hydrocarbons, which improve flame retardance by liberating halogen atoms that retard gas-phase chain oxidation reactions, (b) hydrated metal salts that decompose endothermically in a fire, thus reducing the overall heat of reaction, and which liberate water, which dilutes the flammable gases, and (c) phosphorus-based inorganic and organic additives, which promote the formation of an incombustible char, thus protecting the underlying, unburnt polymer.

It is common practice, especially from a commercial point of view, to use a combination of flame retardants for polymeric materials. In many cases, these flame-retardant mixtures can give an enhanced performance at low cost. The interaction of antimony, most commonly used as antimony oxide, with halogenated polymers or polymers containing halogenated additives, gives rise to a classic case of flame-retardant synergism (Grassie and Scott, 1985). The synergistic effects of phosphorus-nitrogen (Kannan and Kishore, 1992) and phosphorus-halogen (Gou, 1992) are also well-documented. Practical experience has led to the recognition of several combinations of flame-retardant ingredients, and these are frequently employed for flame retarding commercially important plastics.

The alternative method of flame retarding a polymer, namely by chemical modification, has several potential advantages such as: (a) that low levels of modification may suffice; (b) that the modifying groups are chemically attached and therefore less likely to be lost during subsequent service; and (c) that the modification can more readily be molecularly dispersed throughout the polymer.

11.3 Testing procedure and hazard assessments – general aspects

As a consequence of the complex nature and poor reproducibility of fire, there are many techniques for estimating the flammability characteristics of polymeric materials. The most widely used laboratory test is the limiting oxygen index (LOI) measurement, which is a very convenient, precise and reproducible technique (Fenimore and Martin, 1966). The LOI is a measure of the volume percentage of oxygen in a mixed oxygen and nitrogen gas stream that just supports candle-like combustion of a polymer sample. This value therefore enables the combustibility of a polymer to be expressed and compared with that of other materials (Table 11.1). However, the relatively high concentrations of oxygen used in making measurements of LOI are unrepresentative of conditions in a real fire, and generally there is a lack of correlation between the results of most small-scale and full-scale tests.
Medium-scale experiments, based on oxygen consumption calorimetry, such as cone calorimetric measurements, are generally considered to generate parameters that are more relevant to real fire scenarios involving polymeric materials (Babrauskas, 1984). The measured parameters include time to ignition, heat release rate, total heat released, mass loss rate, effective heat of combustion, smoke specific extinction area, smoke production rate and total smoke production (Table 11.2). The central objective in carrying out oxygen consumption calorimetric measurements is to obtain parameters that represent the true nature of the fire hazards posed by the materials under investigation, such as heat release rates (as well as the total heat released) and production of smoke and CO (Schartel et al., 2005) (Table 11.3). The measurements should be carried out on sufficiently thick samples to avoid thermally thin behaviour, and by using a sample holder with minimum heat losses so as to minimize the effects of the apparatus.
employed on the values of the parameters obtained. This will ensure that the values collected through such measurements are in turn more global in nature (de Ris and Khan, 2000).

The burning behaviour of fabrics comprised of a given fibre type or blend is influenced by a number of factors including the nature of the ignition source and time of its impingement, the fabric orientation and point of ignition (e.g. at the edge or face of the fabric or top or bottom), the ambient temperature and relative humidity, the velocity of air, and last but not least on fabric structural variables. Generally, low fabric area density and open structures aggravate the burning rate and thus increase the hazards of burn severity as compared to heavier and multi-layered structures (Backer et al., 1976). In addition, the fabric flammability is determined not only by the fibre behaviour but also by the physical geometry of the fibrous arrays in fabrics. Owing to multi-factorial dependence of the ease of ignition of textile materials, as given by the measured LOI values, several researchers have tried alternative techniques for measurement of the flammability of fabrics to yield values that represent the flammability as an intrinsic property of the material (Miller et al., 1973; Stuetz et al., 1980; Horrocks et al., 1989).

It should be noted here that nearly every country has its own set of standard textile fire testing methods, which are claimed to better represent the special social and technical factors peculiar to each. In addition, test methods are defined by a number of national and international bodies such as air, land and sea transport authorities, insurance organizations, and governmental departments relating to industry, defence and health, in particular. Detailed accounts of different categories of standard test methods advocated for different classes of textile materials are given in detail elsewhere (Horrocks, 2001) (Table 11.4).

Table 11.3 Toxic and asphyxiating gases from the combustion of polymers

<table>
<thead>
<tr>
<th>Gas</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO, CO₂</td>
<td>All organic polymers</td>
</tr>
<tr>
<td>HCN, NO, NO₂, NH₃</td>
<td>Wool, silk, nitrogen-containing polymers</td>
</tr>
<tr>
<td>HCl, HF, HBr</td>
<td>PVC, PTFE, polymers containing halogenated flame retardants</td>
</tr>
<tr>
<td>Alkanes, Alkenes</td>
<td>Polyelefins and other organic polymers</td>
</tr>
<tr>
<td>Benzene</td>
<td>Polystyrene, PVC, aromatic polyesters</td>
</tr>
<tr>
<td>Phenol, Aldehydes</td>
<td>Phenolic resins</td>
</tr>
<tr>
<td>Acrolein</td>
<td>Wood, paper</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>Polyacetals, formaldehyde-based resins</td>
</tr>
<tr>
<td>Formic and acetic acids</td>
<td>Cellulosics</td>
</tr>
</tbody>
</table>
Table 11.4 Selected test methods for textiles

<table>
<thead>
<tr>
<th>Nature of test</th>
<th>Textile type</th>
<th>Standard</th>
<th>Ignition source</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Night wear</td>
<td>BS 5722: 1991</td>
<td>Small flame</td>
</tr>
<tr>
<td>ISO vertical strip similar to tests 1 and 2 in BS 5438</td>
<td>Protective clothing (now withdrawn)</td>
<td>BS 6249: Part 1: 1982</td>
<td>Small flame</td>
</tr>
<tr>
<td></td>
<td>Vertical fabrics</td>
<td>BS EN ISO 6940/1:1995</td>
<td>Small flame</td>
</tr>
<tr>
<td>Small-scale composite test for furnishing fabric/fillings</td>
<td>Furnishing fabrics</td>
<td>BS 5852: Pts 1 and 2: 1979 (retained pending changes in legislation)</td>
<td>Cigarette and simulated match flame (20 s ignition)</td>
</tr>
<tr>
<td></td>
<td>Furnishing fabrics</td>
<td>BS 5852: 1990 (1998) replaces BS 5852: Pt 2</td>
<td>Small flames and wooden cribs applied to small and full scale tests</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ISO 8191: Pts 1 and 2 (same as BS 5852: 1990)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>BS EN 1021-1: 1994</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>BS EN 1021-2: 1994</td>
<td></td>
</tr>
<tr>
<td>Cleansing and wetting procedures for use in flammability tests</td>
<td>All fabrics</td>
<td>BS 5651: 1989</td>
<td>Not applicable but used on fabrics prior to submitting for standard ignition tests</td>
</tr>
<tr>
<td></td>
<td>Commercial laundering</td>
<td>BS EN ISO 10528: 1995</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Domestic laundering</td>
<td>BS EN ISO 12138: 1997</td>
<td></td>
</tr>
</tbody>
</table>

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Air seat assemblies, so called ‘Boeing’ test
All fabrics/composites
ASTM E9060 1983, uses Ohio State University heat release calorimeter
NF P 92501-7, French ‘M test’

Irradiate under \(35 \text{ kW m}^{-2}\) with small burner

Protective clothing
Resistance to radiant heat
Resistance to molten metal splash
Gloves
Fire fighter’s clothing
General flame spread
General protection
Contact heat transmission

<table>
<thead>
<tr>
<th>Nature of test</th>
<th>Textile type</th>
<th>Standard</th>
<th>Ignition source</th>
</tr>
</thead>
</table>
| Use of radiant flux     | Air seat assemblies, so called ‘Boeing’ test
All fabrics/composites | ASTM E9060 1983, uses Ohio State University heat release calorimeter
NF P 92501-7, French ‘M test’ | Irradiate under \(35 \text{ kW m}^{-2}\) with small flame igniter |
| Protective clothing     | Resistance to radiant heat
Resistence to molten metal splash
Gloves
Fire fighter’s clothing
General flame spread
General protection
Contact heat transmission | BS EN 366: 1993) replaces BS 3791: 1970
BS EN 367: 1992
BS EN 373: 1993
BS EN 407: 1994
BS EN 469: 1995
BS EN 532: 1994 (replaces BS 5438)
BS EN 533: 1997 (replaces BS 6249)
BS EN 702: 1994 | Exposure to radiant source
Determine heat transfer index
Molten metal
Radiant, convective and molten metal
Small flame
Small flame
Small flame
Contact temps. 100–500°C |
11.3.1 Specific fire hazards associated with textile fibres

The major fire hazard associated with textile materials, in general, is shown in the high incidence of their implication in fire initiation and propagation and in the related numbers of fatal and non-fatal casualties, especially in domestic environments. Sources of relatively low intensities (such as smokers' materials) are often sufficient to bring about ignition of these materials. Furthermore, conventional textile fibres are comprised of relatively flammable fibre-forming polymers which, owing to their high specific surface area in the fibre form, significantly enhance the rate of combustion and fire proliferation. Coupled with this is the use of textile materials in vertical geometries (e.g. curtains, clothing, etc.) which encourage high burning rates when ignition is at the bottom. Textiles in upholstered furnishings and bedding are also particularly hazardous as is shown in statistics relating to the occurrence of fires in dwellings (Office of the Deputy Prime Minister, 2004). In fact, about 20% of dwelling fires are caused by textile materials (being the first material to be ignited), and about 50% of the fatalities are associated with these fires. Consequently, most flame-retardant textiles are designed to reduce their propensity to undergo ignition and flame propagation when exposed to small flame sources (e.g. burning matches) or to pilots of low intensity but of prolonged duration (e.g. burning cigarette ends). Heat-resistant textiles are not only expected to resist ignition and have reduced flame propagation rates, but also should offer a barrier to heat and flame penetration to the underlying materials or surfaces. Thus, while drapes and curtains may be rendered flame retardant, upholstered furnishing fabrics and protective clothing must be rendered sufficiently flame and heat resistant to prevent damage to underlying fillings, clothing layers or skin (Horrocks et al., 2004).

11.3.2 Flame-retardant strategies for textile materials – general considerations

Textile materials may be rendered flame retardant through a variety of ways. They include: (a) post-chemical treatment of the otherwise flammable component fibres; (b) by the use of fibres that have been flame retarded during manufacture; (c) by the use of inherently flame-retardant fibrous components; (d) or by a suitable combination of the above. For example, conventional cotton and wool fabrics are flame retarded by the use of flame-retardant chemical treatments, which are often simple topical applications of flame-retardant salts (e.g. ammonium polyphosphates), coatings of retardant species (e.g. antimony trioxide/halogenated organic resin systems) or functional finishes (e.g. organophosphorus- and nitrogen-containing monomers, or polycondensates). Functional finishes are
particularly advantageous in that high levels of durability are achieved as they either polymerize within the internal fibre voids or react with functional groups in the fibre-forming polymer backbone (Horrocks, 1996).

The demands of upholstery furnishings to satisfy the aesthetic requirement of the consumer and regulatory flame-resistance standards have seen a significant rise in the use of back-coating treatments, which comprise the flame-retardant species bonded with a resin to the reverse of the fabric. Careful selection of flame retardant and resin can produce a char-generating combination which, when applied to one side (usually the back side) of a flammable textile, confers flame retardancy. This, in the case of thermoplastic fibres, offers a char-forming support, which prevents hole formation. Consequently, back coatings have been successfully applied to fabrics compris of cotton, polyacrylic and polyester blends and mixtures of these fibres. Flame-retardant species can be antimony-bromine combinations, phosphorus-containing species or mixtures of both, often applied with a char-forming acrylic resin.

In the case of common synthetic fibres like polyesters, polyamides (nylons), polyacrylics and polypropylene, it has proved difficult to effectively flame retard owing to: (a) the fact that they do not have physical or chemical structures that are compatible with the more common functional flame retardants; (b) that during fibre production, the melt polymers are often highly reactive and they have low compatibility with many flame retardant additives; (c) that the presence of relatively high flame retardant concentrations (5–20% w/w) necessary to confer flame-retardant properties not only creates spinning fluid compatibility problems but also causes serious reductions in ultimate fibre tensile and other essential textile properties; and (d) that their thermoplasticities, coupled often with a tendency to melt, mean that shrinkage and melt dripping (often under flaming mode) are problems to addressed in the context of secondary fire hazards posed by these materials (Horrocks et al., 2005).

11.4 Polyesters

The linear, thermoplastic, polyesters are all partly crystalline polymers, soluble above their melting points \( (T_m) \) in a range of organic solvents, have glass transition temperatures \( (T_g) \) and therefore softening points that are dependent upon their precise chemical structures, and are highly flammable in their pure state with limiting oxygen indices (LOIs) typically of around 20. The most important polyesters from which useful fibres can be spun are the aromatic polyesters, especially poly(ethylene terephthalate) (PET) and poly(butylene terephthalate) (PBT), owing to their relatively high crystallinities (and hence tensile strengths), \( T_m \)s and \( T_g \)s. However, of the aromatic polyesters, only PET is in wide-spread use as a commercial
flame-retardant polyester and polyamide textiles

PET, when heated to above 300°C, undergoes thermal degradation to give a variety of highly volatile products, many of which are flammable, such as acetaldehyde, methane, ethene and diethyl ether together with less volatile, but none-the-less combustible, products such as PET oligomers (linear and cyclic), terephthalic acid and ethylene glycol (Sobue and Kajiura, 1959; Kardash et al., 1964) The rate-determining step is olefin elimination from glycol dibenzoate groups leading to chain scission and the production of acid and vinyl end groups (Goodings, 1961). The kinetics of this degradation have been thoroughly explored and mechanisms advanced for the production of most products, including a small amount of a carbonaceous residue (Holland and Hay, 2002). Since the pyrolysis of PET gives mainly volatile and flammable products, it would seem probable that vapour-phase flame-retardant strategies would be more effective with PET than strategies aimed solely at increasing char formation, or some other condensed-phase mechanism; however, research to date shows that, in general, mixed phase mechanism can be engineered (Levchik and Weil, 2004).

Progress in the research, development and commercial implementation of flame-retardant strategies for thermoplastic polyesters in general, irrespective of end-application, has recently been comprehensively reviewed by Levchik and Weil (2005). Thus, it is not necessary to review comprehensively this field again; rather it will suffice merely to highlight current commercial practice, significant past and recent research and development, and likely future research directions with regard specifically to the use of polyesters in fibre form.

Choice of a flame retardant for a fibre-forming polymer that is to be melt spun must take into account the possibility that the additive has the potential to interfere with the spinning process, especially if it is an insoluble particulate. Flame retardants also need to be able to withstand laundry cycles if the fibre is to be used in many textile applications. For these reasons, many established inorganic flame-retardant additives and synergists, such as metal hydroxides, antimony trioxide, ammonium and melamine phosphates, and zinc oxides and stannates, are not used in PET fibres. Even if the flame retardant is soluble in the molten polymer, it may be excluded during subsequent crystallization with the possibility of blooming and eventual loss during service. This possibility militates against the use also of many established low molecular weight halogen- and phosphorus-containing flame-retardant compounds. Thus, much research, development and commercial exploitation of flame retardants for PET fibres, especially recently, has concentrated upon the use of flame-retardant, reactive, comonomers. One early example of such a comonomer

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is ethoxylated tetrabromobisphenol A (I), used originally by DuPont in the manufacture of its Dacron 900F® fibre. Such fibres, containing typically up to 6 wt% Br, have LOIs of around 26.

![Chemical structure of ethoxylated tetrabromobisphenol A (I)](image)

Reactive comonomers containing organo-phosphorus groups also have been developed. Thus flame retardancy has been achieved in commercial PET fibres by incorporation of 2-carboxyethyl(methyl)phosphinic acid (II), 2-carboxyethyl(phenyl)phosphinic acid (III) or their cyclic anhydrides. PETs based on the former are marketed under the trade name Trevira CS® (www.treviracs.com). It has been suggested that fibres containing either II or III might be further improved by the additional incorporation of aromatic dicarboxylic acid monomers to act as charring agents (Asrar et al., 1999). Another commercially utilized P-containing comonomer is the 9,10-dihydro-9-oxa-10-phosphaphenanthrenyl-10-oxide (DOPO) adduct of itaconic acid (IV) (Endo et al., 1978). Filament fibres and fabrics based on PET copolymers containing this flame-retardant comonomer are commercially available from Toyobo under the trade name, HEIM® (www.toyobo.co.jp) and have LOIs ranging up to 28.

![Chemical structures of reactive comonomers](image)

It has been reported that the effectiveness of copolymerized DOPO-type monomers can be further improved if alcohohamine derivatives of DOPO, e.g. V, are used rather than similar structures not containing nitrogen (Kim, 1988). Of the FR fibres based on P-containing comonomers, it has been found that those based on IV are the more hydrolytically stable, presumably because the P-containing group is in a side chain rather than
in the main chain (Sato et al., 2000). All the phosphorus-modified PETs, however, appear to be subject to both vapour-phase and condensed-phase mechanisms of flame retardance, with the former predominating (Day et al., 1981; Chang and Chang 1999).

Other reactive comonomers reported to improve flame retardance in PET fibres include phosphinic acid derivatives of itaconic acid (Horie, 2002), and aliphatic and alicyclic (spiro) bisphosphonates such as VI (Murayama and Kashihara, 1978).

![Chemical structures](image)

(V) (VI)

Whilst the reactive incorporation of flame retardants into polyesters may be the optimum approach to achieve effective long-term fire protection, there have been many attempts to flame retard polyester fibres using halogen and/or phosphorus containing additives. Such additives may be applied during the manufacture of the fibres or, more usually, may be applied subsequently to fabrics produced from fibres. An early commercial example of the additive approach was the use of TRIS (tris-(2,3-dibromopropyl) phosphate, (VII)) to flame retard PET fibre in the 1960s and 1970s. There have been several studies over the years of the mode of action of TRIS, between them giving somewhat contradictory indications. However, the emergent consensus seems to be that TRIS acts mainly in the condensed phase with the bromine helping to retain phosphorus species in the condensed phase where they catalyse various condensation reactions, leading to significant char formation (Bostic, Jr. et al., 1973; Inagaki et al., 1977).

Cyclic phosphonates with high phosphorus contents (VIII), manufactured by Rhodia and now marketed by Albemarle as members of their Antiblaze® range of flame retardants, have also been used as additives in flame-retardant PET fibres (www.albemarle.com). They are applied as aqueous solutions to the surfaces of the fibres, which are then heated to soften and swell them, allowing ingress of the additive. The process is similar to the ‘thermosol’ process used in the disperse dying of PET fibres. Mixed vapour-/condensed-phase action has been demonstrated for these types of additive (Day et al., 1981). Cyclic and linear polyphosphazenes also have been suggested as flame-retardant additives for PET fibres (Masaroni and Shintarou, 1998). However, the known instability of such materials to hydrolysis would seem to rule them out for long-term use.
A further type of additive strategy employed with PET is the blending of unmodified PET fibres with flame-retarded fibres or polymer of another type. The use of PET in combination with flame-retarded cotton is well established, but use of other flame-retarded polymers is possible. For example, triphenylphosphine oxide (TPPO) used in conjunction with nylon-6 has been shown to be a good flame retardant for PET fibres. Nylon appears to work synergistically with TPPO by preventing its low temperature sublimation (Bostic and Barker, 1977).

\[
\text{(VII) } \quad \text{(VIII)}
\]

### 11.5 Polyamides

Owing to their aromatic structures and relatively low hydrogen atom contents, the aromatic fibre-forming polyamides, Kevlar® and Nomex® (both developed by DuPont) are naturally flame retardant with LOIs of around 29 and produce high char yields on burning. However, the principal aliphatic fibre-forming polyamides, nylon-6 and nylon-6,6, are less flame resistant: both have LOIs of around 21, depending upon formulation.

Like the linear polyesters, the nylon’s ‘crack’ on heating to give a variety of volatile and flammable products including, in the case of nylon-6, the monomer, caprolactam. Despite this, combustion of polyamides can be relatively slow and small amounts of char may be formed, depending upon the circumstances (Levchik et al., 1999; Levchik and Weil, 2000).

The challenges presented in attempting to flame retard the polyamides are similar to those presented with linear polyesters. That is, any additives or reagents must not substantially interfere with the melt spinning process, adversely affect crystallinity and hence physical and mechanical properties of the fibre, nor be susceptible to degradation and loss during subsequent service, including during any cleaning operations. Thus it is not surprising to discover that rather more progress has been made with flame retarding polyamides for applications in plastic mouldings than for applications as fibres (Weil and Hevehik, 2004). Many of the successful strategies for flame retarding nylon thermoplastic moulding materials, such as use of metal hydrates, inorganic amine phosphates, halogenated and phosphorus-containing organics, and especially red phosphorus, are ruled out for most
applications of nyons as fibres by consideration of effects on melt spinning, fibre properties and subsequent use. As yet, it is not clear that there has been any successful commercial production of flame-retarded nylon textile fibres incorporating flame-retardant additives or reactives, although both Monsanto and Solutia have patented phosphorus-containing diacids (structures IX and X, respectively) designed to replace some of the adipic acid in the manufacture of flame-retardant nylon-6,6 for fibres (Picket and Stoddard, 1977; Asrar, 1988).

\[
\begin{align*}
\text{(IX)} & : & \begin{array}{c}
R - P - \overset{\circ}{\text{COOH}} \\
\text{COOH}
\end{array} \\
\text{(X)} & : & \begin{array}{c}
\text{P} - \overset{\circ}{\text{OH}} \\
\text{COOH}
\end{array}
\end{align*}
\]

However, for non-textile applications of nylon fibres, use of some well-established flame-retardant additives may be possible. Thus Dechloriane Plus\textsuperscript{®} (Occidental Chemical Corporation) (XI) or some other halogenated flame retardant may be added to nyons for use in the melt spinning of carpet fibres (Stoddard et al., 1975).

\[
\begin{array}{c}
\text{(XI)} : \begin{array}{c}
\text{Cl} \\
\text{Cl} \\
\text{Cl} \\
\text{Cl} \\
\text{Cl}
\end{array}
\end{array}
\]

To flame retard nylon textiles and more complex nylon fibre-based articles, there are several commercially established processes whereby treatments are applied externally in manners similar to those employed with polyesters. A well-established finish for nylon textiles is with proprietary Flamegard\textsuperscript{®} (Bayer), thiourea-formaldehyde, resins, which are heat cured with an acid catalyst. Chemical grafting of materials to the surfaces of fibres has also been tried. For example, nylon-6 fibres have been surface grafted with poly(2-methyl-5-vinyl pyridine) and then further treated with various halogenated flame retardants to give LOIs of up to 31 (Efros et al., 1983) and with o-chlorophenol to similar effect (Mukherjee et al., 1981).
11.6 Conclusions and future trends

It seems unlikely that there will be any major breakthroughs with regard to new and/or improved reactive flame-retardant comonomers or conventional organic and inorganic flame-retardant additives for use in either PET or nylon fibres. The requirements to achieve satisfactory flame retardance without appreciably interfering with the spinning process, modifying the physical and mechanical properties of the fibres, or affecting long-term stability, and at economic cost, restricts options considerably.

However, since the important first barrier to ignition of a fibre (or indeed of any material) is the surface, it is sensible to ask whether or not more can be done to modify fibres or textile surfaces so as to improve flame retardance by surface modification, especially using modern vacuum or plasma deposition techniques. Some encouraging results have been reported on the plasma modification of the surfaces of polyacrylonitrile fibres so as to improve flame retardance (Akovali and Gundogan, 1990), which suggest that the technique might be more universally applicable. Already, plasma modifications of polyester fibres and fabrics have been reported aimed at modifying surface adhesion and reflectivity (Manenq et al., 1999; Van Ooij et al., 1999; Lee et al., 2001). Conventional vacuum coating might also be appropriate for the deposition of some types of flame-retardant surface modification to fibres and fabrics (Mikael and Yializis, 2005).

An alternative type of flame-retardant surface treatment for fibres and fabrics is the intumescent coating, applied using conventional coatings technology. Horrocks and colleagues have achieved considerable success with this type of approach, especially for the flame retardation of cotton and wool fabrics, but the methodology is, in principle, extendable to other types of polymer-based textile (Horrocks, 1996).

Nanoscopic materials are attracting increasing interest, both for surface coatings and as additives in polymers. The Nano-Tex® (www.nano-tex.com) surface-treatments for fabrics, based on dispersions of nanoscopic additives, and designed thus far to improve stain resistance, static build-up, and moisture dissipation in fabrics, would appear to be extendable to the deposition of nanoscopic flame-retardant formulations. The exploitation of nanoscopic particles within polymer-based fibres is also growing rapidly with the aim being mainly to improve modulus and, if possible, also tensile strength. Fibres have recently been spun successfully from polyesters containing, for example, nanoscopic organoclays (Chang et al., 2004; Mun et al., 2006), carbon nanotubes (Yang et al., 2006) and fullerenes (Ikegame and Kono, 2006). Since the incorporation of selected nanoscopic materials into bulk polymers, along with conventional flame retardants, can significantly improve their flame retardancy, this is clearly an area that is ripe for exploitation with fibres. It is to be noted that some success with this
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11.7 Sources for further information and advice


11.8 References


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Flame-retardant polyester and polyamide textiles


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12 Advances in functional finishes for polyester and polyamide-based textiles

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12.1 Introduction

12.1.1 Nylon

Nylon was the first truly synthetic fibre which at the time of its introduction in the late 1930s was supposed to have the potential to replace many of the natural fibres. It had strength, lustre, softness, recovery, resilience, good abrasion resistance, crease recovery, easy dyeability, etc. on the positive side. Eventually it did not do so, and it was primarily due to its plastic feel and shine in flat form, poor moisture regain and the propensity to develop static charge.

The initial use for the nylon filaments was hosiery, especially stockings due to its excellent recovery and shape retention properties. Most of the nylon was used in military applications especially parachutes during the World War II, which shortly followed the development of nylon. Initially an attempt was made to address the problem of durability of the apparels made from wholly natural fibres by blending nylon with cotton. However nylon could never really penetrate the woven textile segment and the introduction of superior PET/cotton blends stopped any further scope for nylon in this sector. Since then nylon has remained confined to some specific application areas which exploit its properties to the best.

12.1.2 Polyester

Polyester, which followed nylon four years later, today commands a lion’s share of synthetic textile fibres. The positive attributes of polyester fibres are strength, wrinkle resistance, good wash and wear properties, durability, etc. Although it has poor moisture regain, this has been partly countered by blending it with natural/man-made fibres having higher moisture regain. In fact, polyester-cotton and polyester viscose blends today are the most important textile types after 100% cotton textiles in terms of volume generated annually worldwide.
**Table 12.1 Tensile properties of nylon and polyester fibres**

<table>
<thead>
<tr>
<th>Property</th>
<th>Nylon 6,6</th>
<th>Nylon 6</th>
<th>Polyester</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tenacity ($N,\text{tex}^{-1}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>0.4–0.6</td>
<td>0.4–0.6</td>
<td>0.4–0.6</td>
</tr>
<tr>
<td>High tenacity</td>
<td>0.6–0.9</td>
<td>0.6–0.9</td>
<td>0.8–1.0</td>
</tr>
<tr>
<td>Breaking extension (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>20–30</td>
<td>20–40</td>
<td>10–20</td>
</tr>
<tr>
<td>High tenacity</td>
<td>15–20</td>
<td>15–20</td>
<td>5–10</td>
</tr>
<tr>
<td>Initial modulus ($N,\text{tex}^{-1}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>2.0–3.5</td>
<td>1.5–3.5</td>
<td>8–10</td>
</tr>
<tr>
<td>High tenacity</td>
<td>4.0</td>
<td>5.0</td>
<td>4.0</td>
</tr>
<tr>
<td>Work of rupture ($\text{mN,\text{tex}^{-1}}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>60–70</td>
<td>70–80</td>
<td>50–60</td>
</tr>
<tr>
<td>High tenacity</td>
<td>50–60</td>
<td>60–70</td>
<td>20–30</td>
</tr>
</tbody>
</table>

**Table 12.2 Typical values for the yield strain and elastic recovery of various fibres**

<table>
<thead>
<tr>
<th>Material</th>
<th>Yield strain (%)</th>
<th>Elastic recovery % from</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1% extension</td>
</tr>
<tr>
<td>Nylon</td>
<td>16</td>
<td>90</td>
</tr>
<tr>
<td>Polyester</td>
<td>3</td>
<td>98</td>
</tr>
<tr>
<td>Cotton</td>
<td>1</td>
<td>91</td>
</tr>
</tbody>
</table>

**12.2 Properties and uses of polyester and polyamide fibre/textiles**

Unlike the natural fibres the properties of synthetic fibres depend to a large extent on the morphology of the fibre, which in turn is governed by the molecular weight of polymer, extrusion and drawing conditions, and the thermal history of the fibres. Tables 12.1 and 12.2 give the general tensile and the recovery properties of nylon and polyester fibres.¹

What distinguishes the synthetic fibres from natural fibres is their high strength, durability, better chemical resistance, uniformity, crease recovery and ability to be tailored to different mechanical properties by suitable modification of fibre formation parameters.

To a large extent the application areas for any fibrous material are determined by its mechanical properties. A look at Tables 12.1 and 12.2 makes it obvious that the yield and recovery behaviour of polyester and cotton are similar. Polyester also has a much higher initial modulus than polyam-
ide and similar to that of cotton. Hence in a polyester-cotton blended yarn, the contribution of the two components to the overall strength is better as the stress–strain curves of cotton and polyester are similar. However, due to the lower initial modulus of polyamide fibres, the contribution of the polyamide component to the overall strength in a cotton-polyamide blend is poor. This has resulted in huge popularity and success of polyester-cotton blends. Polyamide, on the other hand has excellent recovery properties which make it ideal material for knitted garments where better fit and shape retention are more important. However, polyamide is a versatile fibre that can be used in a wide range of applications: fashion garments, functional sports and leisurewear, ladies’ hosiery, lingerie, sewing thread, carpets, upholstery and even technical applications such as balloons and parachutes, sails, climbers’ ropes and automotive fabrics. It is also increasingly being used in blends with natural fibres and other man made fibres.

Polyamide fibres are available in a wide range of linear density and cross-sections: ultra-fine microfibres of less than 1.0 dtex for lightweight articles such as hosiery and lingerie to coarse fibres of up to 2100 dtex for robust articles such as suitcases, rucksacks and shoes. Round, trilobal and dumb-bell shaped cross-sections are most common. Number of filaments in the yarn varies enormously and it is this that determines the properties of the end product.

12.3 Imparting functionality through finishing/coating/laminating

Although synthetic fibres like polyester and polyamide have many positive attributes, their inherently low moisture regains and high strength result in problems of static generation, easy soiling and pilling. Part of these problems can be overcome by blending these fibres with natural fibres. However, in applications where synthetic fibre component is very high (or 100%), these problems remain. Additionally, there are greater functional demands from these fibres. On one hand, the ever more demanding customer requirements require the traditional apparel garments to be antistatic, antimicrobial, easy to clean, stain proof, etc. On the other hand the increasing use of these fibres for technical applications demands functionalities like water-repellency, water-proof breathability, gas barrier, flame-proofness, UV protection to name a few from textiles made from these fibres.

There are basically four approaches to impart functionality to the textiles made from polyester and polyamide textiles:

- By incorporating a suitable additive in the melt or dope of the fibre polymer to produce intrinsically functional filaments (e.g., Trevira, the intrinsically flame retardant polyester fibre)
• Applying a suitable chemical/mechanical finish to the textile
• Applying a coating of a functional chemical formulation on the textile substrate
• Laminating the textile with a preformed film or layer (an adhesive would be needed to bond the film/layer with the textile).

It must be noted here that coating and lamination approach to impart functionality would alter many of the desired properties of an apparel textile like flexibility, breathability, drape, etc. and hence may not be suitable for such applications. Discussions on imparting functionality at melt or dope stage is beyond the scope of this topic and hence only the remaining three approaches would be discussed.

12.3.1 Antistatic

One of the problems generally faced while using wholly or predominantly synthetic textiles is their tendency to generate static electricity. In most cases the generation of static charge is a minor irritant leading to small problems like clinging of garments, mild shocks, increased soiling, but in rare cases can lead to fire, explosion and damage to electronic components. The problem of static generation becomes acute with synthetic fibres in low humidity conditions. The static generation can be reduced by increasing the humidity of the workplace, incorporating the antistatic agents in the manufacture of the fabric. A detailed discussion on the causes and solution of static generation problem can be found elsewhere.

The static generation may manifest itself in problems at two distinct stages. There may be problems related with the production of textiles, i.e.,

1. repulsion of fibres from each other in charged slivers
2. lapping of fibres around rollers
3. ballooning of yarns
4. electrostatic attraction between fabrics and lint/soil.

Then there may be problems related to the use of the textiles, like generation of static on walking over a carpet, etc. Majority of the chemical finishes used as the antistatic agents for synthetic fibres today are the variants of the finishes which have been in the market since the 1970s. Traditionally the antistatic finishes for polyamides have been based on ammonium salts, quaternary ethoxylated amines and quaternary fatty amide amines.

The static problem on polyester can be reduced to some extent by sodium hydroxide treatment (weight reduction). The effectiveness of the treatment
can be increased by finishing it with a reactive polyethylene glycol of mol. wt 1000 which is durable up to 30 launderings. Silane nanosol has been successfully prepared with an organic precursor to improve the anti-static property of polyester fabrics. The experimental results show that the anti-static property of treated PET fabrics was enhanced considerably by sol treatment, while there were no negative effects on the whiteness, strength and handle of the treated fabrics. Static charge usually builds up in synthetic fibres such as nylon and polyester because they absorb little water. Cellulosic fibres have higher moisture content to carry away static charges, so that no static charge will accumulate. As synthetic fibres have poor anti-static properties, research work concerning the improvement of the anti-static properties of textiles by using nanotechnology has been undertaken. It was determined that nano-sized titanium dioxide, zinc oxide whiskers, nano antimony-doped tin oxide (ATO) and silane nanosol could impart anti-static effects because they are electrically conductive materials. Such materials help to effectively dissipate the static charge which is accumulated on the fabric. On the other hand, silane nanosol improves anti-static properties, as the silane gel particles on fibre absorb water and moisture in the air by amino and hydroxyl groups. Nanotechnology has also been applied in manufacturing an anti-static garment. W. L. Gore and Associates GmbH used nanotechnology and polytetrafluoroethylene (PTFE, Dupont’s Teflon®) to develop an anti-static membrane for protective clothing. Gore-Tex® I Workwear protects the wearer from electrostatic discharges. Electrically conductive nano-particles are durably anchored in the fibrils of the Gore-Tex® I membrane of Teflon, creating an electrically conductive network that prevents the formation of isolated chargeable areas and voltage peaks commonly found in conventional anti-static materials. This method can overcome the limitation of conventional methods, which is that the anti-static agent is easily washed off after a few laundry cycles.

12.3.2 Antimicrobial

There are two distinct roles the antimicrobial finishes have to play: firstly the prevention of the growth of disease causing microorganisms on textiles which may not cause biodeterioration of textiles but can cause malodour, stains and cross-infections. Second, the protection of the textile itself from attack by the mildew, mould and rot producing microorganisms. The synthetic fibres especially polyester and nylon are generally not attacked by microorganisms; hence the discussion on antimicrobial finishes in this section will be confined to the finishes which deal with disease and odour causing microorganisms.
An ideal antimicrobial for textiles would have to fulfil the following basic requirements. It should provide safety in the form of low toxicity to the consumer, e.g. it would not cause allergy or irritation to the skin. Its application would not adversely affect textile properties or appearance. It should be compatible with common textile processing and the resulting antimicrobial efficacy has to be durable against repeated laundering. Today such finishes are coming under increasingly heavy pressure from regulatory bodies to conform to strict environmental and safety norms.9

There was a spurt in the use of antimicrobial finishes for textiles during World War II. A large number of compounds belonging to many classes like inorganic salts, organometallic compounds, phenols, thiophenols, antibiotics, formaldehyde derivatives, amines, etc. were tried and processes developed to make textiles resistant to microbes.10 Today many of these compounds have been banned or phased out due to their high toxicity and potential to cause environmental damage. Examples of such compounds are copper naphthenate, copper 8-quinolinate and many organo mercury compounds.11 The antimicrobial finish may act by one of three mechanisms; controlled release (chemical or physical), regeneration and barrier layer. The application of the finish itself may be by insolubilization on textile fibre, chemical modification of the fibre by covalent bond formation, coating on the textile fibre surface or microencapsulation.

Some of the old products which still find use are tributyl tin, dichlorophene, 3-iodopropynyl butyl carbamate, benzimidazol, derivatives of salicylanilides and alkylolamide salts of undecylenic acid. The structures of two antimicrobial agents are given in Figure 12.1. A popular and widely used agent is 2,4,4′-trichloro-2′-hydroxyphenyl ether or triclosan. It is used in mouthwashes, toothpastes, liquid hand soaps, deodorant products and also in textiles. Tinosan AM 100 and Tinosan AM 110 are antimicrobial finishing agents based on Triclosan produced by Ciba Specialty

![Molecular structures of two antimicrobial agents.](image)
Chemical Ltd Germany. These products are designed to impart durable antimicrobial efficacy to fabric made by polyester, polyamide fibres and blends of such fibres with cotton and wool. The active antimicrobial agent in these products acts like disperse dye and can diffuse into the fibre with a very high exhaustion rate. During use, the antimicrobial active agent can migrate to the surfaces of the treated textiles at a slow yet sustained rate providing excellent and durable antimicrobial efficacy.  

Durable and regenerable antibacterial textiles were prepared by Sun and coworkers \(^{13}\) by continuous grafting of cyclic amine monomer 3-allyl-5,5-dimethyl hydantoin (ADMH). The textiles selected were nylon, polyester, acrylic, polypropylene and natural fibres. After the grafting process, the treated textiles were exposed to dilute chlorine solutions which convert hydantoin structures to N-halamines, which act as powerful, durable and regenerable antibacterial agents against bacteria \(E\ coli\).  

In a novel process\(^ {14}\) light activated antibacterial nylon textiles were prepared by Bozja by grafting on proporphyrin IX and zinc proporphyrin IX. The fibres were active against \(S.\ aureus\) after exposure to 10000 lux and against \(E.\ coli\) after exposure to 60000 lux. No activity was shown in absence of light.  

A simple way of creating antibacterial nylon fibres has been reported by Sun et al.\(^ {15}\) The carboxylic end groups in nylon were utilized for interaction with cationic quaternary ammonium salts in alkaline conditions. The role of pH, time and temperature and the structure of cationic salt were critical in determining the effectiveness of the process. Use of optimized conditions produced wash fast antibacterial textiles. Using similar strategy, Kim et al. produced antimicrobial nylon 6,6 textiles by treating it with Berberine,\(^ {16}\) a natural cationic colorant.  

Recently cyclodextrins have been finding increasing use in textile finishing due to their ability to form inclusion complexes (ICs) with certain compounds (host) and the property to release the host at a controlled rate. This approach was used by Gawish et al.\(^ {17}\) They grafted glycylid methacrylate (GMA) onto nylon 6 and then reacted \(\beta\)-cyclodextrin (CD) or monochlorotriazine \(\beta\)-cyclodextrin with GMA. The CD cavity was used for the formation of inclusion complexes with some biocidal guests including p-hydroxy benzoic acid, \(\text{AgNO}_3\)-ethanol amine mixture, iodine, N,N-dimethyl tolunamide (DETA), citronella, jasmine and sweet basil. The ICs thus formed act by controlled release of biocide and are active over long periods of time.  

For imparting anti-bacterial properties, nano-sized silver,\(^ {18}\) titanium dioxide and zinc oxide\(^ {19}\) are also used. Metallic ions and metallic compounds display a certain degree of sterilizing effect. It is considered that part of the oxygen in the air or water is turned into active oxygen by means of catalysis with the metallic ion, thereby dissolving the organic substance.
to create a sterilizing effect. With the use of nano-sized particles, the number of particles per unit area is increased, and thus anti-bacterial effects can be maximized. Nano-silver particles have an extremely large relative surface area, thus increasing their contact with bacteria or fungi, and vastly improving their bactericidal and fungicidal effectiveness. Nano-silver is very reactive with proteins. When contacting bacteria and fungus, it will adversely affect cellular metabolism and inhibit cell growth. It also suppresses respiration, the basal metabolism of the electron transfer system, and the transport of the substrate into the microbial cell membrane. Furthermore, it inhibits the multiplication and growth of those bacteria and fungi which cause infection, odour, itchiness and sores. Hence, nano-silver particles are widely applied to socks in order to prohibit the growth of bacteria. In addition, nano-silver can be applied to a range of other healthcare products such as dressings for burns, scald, skin donor and recipient sites.\textsuperscript{20} Titanium dioxide is a photocatalyst; once it is illuminated by light with energy higher than its band gaps, the electrons in TiO\textsubscript{2} will jump from the valence band to the conduction band, and the electron (e\textsuperscript{−}) and electric hole (H\textsuperscript{+}) pairs will form on the surface of the photocatalyst. The negative electrons and oxygen will combine into O\textsubscript{2}−; the positive electric holes and water will generate hydroxyl radicals. Since both are unstable chemical substances, when the organic compound falls on the surface of the photocatalyst it will combine with O\textsubscript{2}− and OH− respectively, and turn into carbon dioxide (CO\textsubscript{2}) and water (H\textsubscript{2}O). This cascade reaction is called ‘oxidation-reduction’, and the mechanism is shown in Figure 12.2. Through the reaction, the photocatalyst is able to decompose common organic matters in the air such as odour molecules, bacteria and viruses. Several

\begin{center}
\includegraphics[width=\textwidth]{photocatalysis.png}
\end{center}

\textit{12.2 Photocatalysis mechanism of titanium dioxide.}
papers have discussed the use of the photocatalytic property of TiO$_2$ in the field of textiles.\textsuperscript{10} It was determined that a fabric treated with nano-TiO$_2$ could provide effective protection against bacteria and the discoloration of stains, due to the photocatalytic activity of nano-TiO$_2$. On the other hand, zinc oxide is also a photocatalyst, and the photocatalysis mechanism is similar to that of titanium dioxide; only the band gap (ZnO: 3.37 eV, TiO$_2$: 3.2 eV) is different from titanium dioxide. Nano-ZnO provides effective photocatalytic properties once it is illuminated by light, and hence it is employed to impart anti-bacterial properties to textiles.\textsuperscript{21}

12.3.3 Soil release/stain proof

Soiling of textiles is a natural process as it increases the entropy of the soil-textile system. The process of soiling may occur mainly by one of three ways, i.e., a direct contact between a textile and soil, a soiled surface coming in contact with a non-soiled one and by electrostatic attraction of airborne soils onto the electrostatically charged textile surfaces.\textsuperscript{22} Soil may be of many types: 1. water soluble; 2. particulate; 3. bleachable; 4. oily or greasy (hydrophobic); 5. proteinaceous; and 6. mixed or composite.\textsuperscript{23}

The adsorption of the soil generally occurs by Vander der Waals forces, which operate only over very short distances. It is the oily or greasy soil that is the most difficult to remove from hydrophobic textile fabrics like polyester, acrylic and nylon due to the difficulty in the wetting of soil-fibre interface. It has been shown that the removal of oily soil involves three stages:

1. An induction period during which wetting of the soil-fibre interface takes place
2. Release of the soil from the textile surface by roll up mechanism and
3. The final stage during which little soil removal takes place.

The rate of soil removal is governed by the length of the induction period which may be short or very long, as wetting of hydrophobic soil-hydrophobic textile system may be slow (Figure 12.3).

The rate and extent of soil removal can be increased by application of an essentially hydrophilic soil release finish on the hydrophobic fibre. This soil release finish facilitates the faster hydration of the fibre surface. The water can diffuse through the soil release film under the oily soil and hasten its removal (Figure 12.4).

To increase the durability of such a soil release finish on a hydrophobic fibre, it must be amphiphilic in nature. The oleophilic part can be anchored on the fibre while the hydrophilic portion is oriented outward. The soil release finish may be classified based on the nature of hydrophilic component of the finish, i.e., carboxylic, hydroxyl or oxyethylene groups. The
Non-polymer treatments

The soil release property of the polyester fabric can also be increased by making the fibre surface hydrophilic by alkaline hydrolysis. Treatment of polyester fabric with 10% NaOH causes formation of –OH and –COOH
groups on fabric surface leading to excellent soil release properties comparable to those obtained with conventional soil release finishes but more durable.\textsuperscript{24}

\textit{Plasma treatment}

Plasma irradiation of nylon and polyester fabric surface is another effective way of improving the surface hydrophilicity. This approach is increasingly being explored for surface modification of textile fabrics. Plasma treatment can be carried out in mainly two ways. In presence of a suitable precursor, a thin polymeric film can be deposited on the fabric surface. The properties of the treated textiles are governed by the chemical nature and the thickness of the deposited film. In absence of any precursor, surface functionalization (by oxidation or hydrolysis) or increase in surface roughness may occur. Remarkable increase in the wettability of polyester and polyamide fabrics was observed\textsuperscript{25–26} on exposure to low temperature plasma. While Okten and Seventekin\textsuperscript{25} reported improvement in soil resistance of the treated fabrics, Kartick \textit{et al.}\textsuperscript{26} report formation of nanosized vertical and horizontal channels on polyester and nylon fabric surfaces. They surmise that these channels may contribute towards increased hydrophilicity of the textiles. Cireli \textit{et al.}\textsuperscript{27} used acrylic acid as the precursor and the reported significant improvement in the wettability of polyester and nylon fabric probably due to the deposition of a hydrophilic polymeric film on the treated fabrics. However Stefecka \textit{et al.}\textsuperscript{28} observed formation of –COOH groups on polyester monofilaments on exposure to atmospheric nitrogen plasma but did not find any evidence of any surface roughness.

12.3.4 Water repellent

A detailed account of the principles and the chemistry of water repellency and water proofing is given by Holme\textsuperscript{29} and Kissa.\textsuperscript{30} Conventionally the approach to water repellency has been the application of a coating of relatively hydrophobic materials like insoluble metal salts of soaps, waxes, organometallic complexes, silicone compounds and lately fluorochemical compounds on the textile substrate. These finishes are usually applied by conventional method of pad-dry-cure. To be effective, the fabric construction needs to close and the finish must be able to coat all the fibres and fill the interstitial gaps between the yarns.

Today, the textile finishes are moving from simple water repellent finishes to ultrahydrophobic/superhydrophobic finishes which are characterized by contact angles in excess of 150°. These are discussed in detail in Chapter 13.
12.3.5 Waterproof/breathable

Waterproof/breathable fabrics are defined as fabrics that will withstand over 1000 mm of water (9.8 kPa) pressure without leaking, whilst allowing water vapour to pass through. In applications like rainwear, outdoor sports clothing, small tents, climbing and mountaineering, etc. Water vapour breathability becomes an important functional requirement. Such textiles are created by coating or laminating some waterproof/breathable material onto a textile substrate.

The current account of waterproof breathable textiles will outline the basics only. For the detailed accounts of the same, the reader is directed elsewhere.\textsuperscript{31–32} The requirement of this particular functionality in waterproof fabrics is needed as the human body produces heat and perspiration at varying rates at almost all times as a result of metabolism. The rate of heat and moisture generation depends on the intensity of the physical activity undertaken. Table 12.3\textsuperscript{31} gives the heat and perspiration produced by the average human body.

According to Table 12.3, human body produces heat and perspiration even when there is apparently no physical activity (sleep). It is important to maintain the body core temperature to feel comfortable. If a person becomes wet in very cold weather, then the core temperature of the body may go down dangerously, resulting in a condition known as hypothermia. On the other hand if the body heat and perspiration are not allowed to escape at a rate sufficient to maintain the body core temperature, then core temperature of the body may rise dangerously, resulting in a condition called hyperthermia. In normal cases, these can cause discomfort, but in extreme cases can lead to injury, incapacitation or even death. Hence for all the apparel applications, which have been given waterproof treatments, breathability becomes a necessary functionality.

\begin{table}[h]
\centering
\caption{Heat energy produced by various activities and corresponding perspiration rates}
\begin{tabular}{lll}
\hline
Activity & Work rate (Watts) & Perspiration rate (gram per day) \\
\hline
Sleeping & 60 & 2280 \\
Sitting & 100 & 3800 \\
Gentle walking & 200 & 7600 \\
Active walking & 300 & 11500 \\
With light pack & 400 & 15200 \\
With heavy pack & 500 & 19000 \\
Mountain walking with heavy pack & 600–800 & 22800–30400 \\
Maximum work rate & 1000–1200 & 38000–45600 \\
\hline
\end{tabular}
\end{table}
are essentially four ways in which the textiles can be made waterproof/breathable:

- densely woven fabrics
- membranes
- coatings and
- biomimetics.

**Densely woven fabrics**

These were first developed during World War II. Very densely woven cotton fabrics were prepared by keeping the fibres as parallel to the yarn direction as possible. In contact with water, the fibres swell to block all pores to prevent the penetration of the fabric by water. The products based on such an approach are still commercially important, Ventile being one of the brand names.

**Membranes**

These are very thin (≈10 µm) polymeric films having a very high resistance to water penetration. These are of two types.

**Microporous membranes**

These membranes are typically made from PTFE (poly tetra fluoroethylene), PVDF (poly vinylidene fluoride) or hydrophobic PU (polyurethane) and have a large number of micropores (1.4 billion/cm²). The pore size is 2–3 µm which is too small to allow water droplets (100 µm) to pass through but large enough for the water vapour to transmit through. Additionally, the hydrophobic nature of the polymer enhances the waterproof effect, although the same may cause the problem of adhesion with the base fabric. One of the most widely known products used in waterproof breathable industry is Gore-Tex® membrane, based on expanded PTFE. Figure 12.5 shows the SEM of a Gore-Tex membrane and its mechanism.

**Hydrophilic membranes**

These are solid monolithic films without any pores. These are prepared from chemically modified polyester or polyurethane. The transmission of the water vapour takes place by diffusion through the membrane. The polyester or polyurethane is modified to incorporate up to 40% polyethylene oxide or polyether groups. These groups form the amorphous part of the membranes and provide the required hydrophilicity for the diffusion of the water vapour.
SympaTex® brand of fabrics in an example of waterproof/breathable fabrics based on copolymer of polyester and polyether. Recently the engineering plastics division of DSM, Sittard/Netherlands has introduced Arnitel polymer. This co-poly-ether-ester based polymer can be made into monolithic films for waterproof/breathable textile applications. Another waterproof/breathable fabric based on similar mechanism is the eVENT brand of fabrics.

**Coatings**

Coatings are also layers of polymeric materials which are much thicker than the membranes. These are also microporous and hydrophilic in nature. The basic chemical nature and the mechanism governing the passage of water vapour in coatings are essentially similar to those of the membranes. However, the rate of moisture transmission decreases sharply with increase in the thickness of the coating.

**Biomimetics**

Biomimetics mimic the biological structures and mechanisms in man-made products. For example, the mechanism of a leaf stomata, which opens when plant needs to transpire more water vapour and closes (Figure 12.6) when it needs to reduce it, has been the inspiration for some products.

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Akzo Nobel has developed one such product called Stomatex®. Stomatex is generally made from thermo-insulating closed-cell foam materials such as Neoprene. It uses a pattern of dome-shaped vapour chambers, each with a tiny pore in the centre. While resting, excess body heat and perspiration rise into the dome-shaped chambers and exit through the tiny pores at a controlled rate. The chambers flex and stretch with each movement of the body, causing excess heat and perspiration to be pumped out of the small pores. This unique action of the vapour chambers increases and decreases with the user’s level of physical activity.

There are other fabric constructions which do not use membranes, coatings or laminates but can keep the wearer dry by directing the water as well as moisture away from the body. Hence these fabrics have a directional effect as far as the transport of water in liquid and vapour forms is concerned. These can also be called breathable. Nikwax Directional Textiles as used by Páramo Directional Clothing Systems and Feather and Fur Technology supplied by FurTech are some examples of such directional fabrics. Links to the brand products of other waterproof/breathable textiles can be found on the internet.

In a breakthrough development, a smart temperature controllable water vapour permeable polyurethane (PU) film has been developed. The film can control its water vapour permeability (WVP) with temperature change by itself. This was achieved by optimizing the hard to soft segment ratio, PEG content and isocyanate index of the PU polymer. It was found that smart PU film is obtained at MDI/PEG/PBA/1,4-BD ratio of 3:0.6:0.4:2. The polymer has a Tg of ~24°C and the WVP starts to increase above 18°C, which is close to the Tg of the PU polymer.
12.3.6 Hydrophilic finishes

Textile fibres with inherently low moisture regain like polyester, nylon, acrylic, polypropylene, etc. can cause problems of static charge accumulation, soil pickup/poor soil release and poor wettability. In many applications improved wettability or moisture absorption is important. There have been attempts to render these fibres hydrophilic to improve their moisture absorption capacity. Many of the soil release finishes are in a sense hydrophilic finishes as they rely on the improved hydrophilic character of the fibre to release hydrophobic oily and greasy soils.

One approach in this direction is to create polar and ionic groups on the surface of the fibres by hydrolysis and plasma treatment. This approach has already been discussed. The other approach is to apply hydrophilic polymers onto the fibre surface and create some kind of bonding between the finish and the fibre to obtain a reasonable degree of finish durability.

In one such approach Roos et al.\textsuperscript{39} used cationic starch, cationic cellulose derivatives and hydrophobically modified cationic cellulose for modification of polyester fabric. The main reason for the adsorption of the cationic polymer on PET surface was slightly negatively charged surface of polyester. Application of the cationic polymers resulted in significant improvement in the wetting behaviour of polyester fabric.

Recently there has been a tendency to apply natural hydrophilic proteins to improve the hydrophilicity of synthetic fibres. Silk sericin, a byproduct of silk degumming process, has received special attention lately in this regard. In one such attempt,\textsuperscript{40} polyester swollen with benzyl alcohol, was grafted with N-vinyl formamide by electron beam irradiation technique. The grafting degree on polyester could be controlled by the swelling degree and by monomer concentration. This was followed by hydrolysis with \textit{H}_2\textit{SO}_4 and sericin was fixed with a cross linking agent, ethylene glycol diglycidyl ether. The grafted fabric showed good wettability and antistatic property. The increase in wettability was due to incorporation of amide groups on the surface of the PET, which was expected also to have antibacterial effect.

Kongdee et al.\textsuperscript{41} used supercritical carbon dioxide to impregnate sericin into polyester. They found no impregnation of PET with sericin if polyester surface is not modified as there is no interaction between hydrophilic sericin and hydrophobic polyester. However, if the PET surface is modified with alkali to generate –OH and –COOH groups on fibre surface, sericin is bound to the polyester fibre surface rendering it hydrophilic.

12.3.7 UV protective finish

Sunlight is an essential source of all life on earth. Although the solar spectrum may extend from below 200 nm to above 3000 nm at sea level, it is...
the lower wavelength portion (200–400 nm), which is important. This is essentially the high-energy portion and generally known as Ultra Violet radiation (UVR). Whereas the visible region of the solar spectrum is essential for most life sustaining processes on earth, it is the UVR that can cause some undesirable effects due to its high energy.\(^4\)

Almost all the radiation of wavelength below 290 nm is filtered out by the ozone layer in earth's atmosphere. The UV part of the solar radiation (290–400 nm) is beneficial to human beings in small doses; however, large doses, especially in the short UVB range (280–315 nm), may cause sunburns, skin cancer, photokeratitis, photodermatosis, etc.\(^43\)–\(^44\) In addition to its deleterious effects on human beings, it also deteriorates the material properties of apparel, upholstery, draperies, carpets, furniture, paints, electronic parts, building construction materials – wood, plastic panels, etc. and other articles of outdoor use and limits their durability and life span.

With the alarming increase in the rate of ozone layer depletion in the earth’s atmosphere, the risks involved due to prolonged exposure to solar UV radiation are increasing day by day. Hence it becomes imperative to protect the human skin and other materials from harmful effects of solar UV radiation. There are various approaches to tackle this problem. For protection of human skin, use of sunscreen lotions and other cosmetics is quite common. The experts have recommended use of textiles capable of blocking harmful UV radiation during outdoor activity. Recently much work has been done to develop special UV protective clothing and consideration has been given to factors affecting UV protection capability of fabrics like fibre type, cover factor, construction (weave/knit pattern), porosity, thickness, presence of dyes/pigments,\(^44\)–\(^45\) etc. The ability of a particular textile to block UV radiation is measured in terms of SPF (sun protection factor) or UPF (ultra violet protection factor), which depends on the fibre type and various other fabric construction parameters. A value of 30 and above is supposed to provide good protection. Special UV protecting chemicals and processes have been developed\(^46\)–\(^48\) that enhance the UV protective capability of the textiles. Ciba-Geigy has launched a number of UV absorbers for different textile fibres, which are available in the brand names like Cibatex W, Cibatex APS, Ciba Tinofast CEL, Ciba Tinofast PES, etc. and can be applied by exhaust or pad-batch methods. Similarly, a process developed by Textile Department of the University of New South Wales and launched by Clariant is known as Rayosan process.\(^49\) The two products available are Rayosan C Paste and Rayosan CO Liquid. In this technology fibre reactive Rayosan products can be applied on cellulosic, wool and nylon fibres during dyeing.

The active chemical responsible for the UV protection should be able to absorb solar UV radiation efficiently (esp. in high energy UVB region,
280–320 nm), convert the absorbed energy into vibrational energy and then into heat energy to the surroundings without causing photodegradation of the fibre or the finish itself. Structures of some common UV absorbers are given in Figure 12.7.

Some interesting UV resistant fibres have been developed with a sheath core structure wherein the core consists of polyester, polyether ester, polyester urethane, polyester amide,\textsuperscript{50} polyamide,\textsuperscript{51} polyester with 10\% TiO\textsubscript{2}, ZnO and/or alumina\textsuperscript{52} and the sheath consists of similar polymers with UV absorbers i.e., carbon black, etc. Similarly titanium dioxide (TiO\textsubscript{2}) that is sometimes incorporated in man-made fibres for delustring, also provides protection from UV radiation.

Inorganic UV blockers are preferable to organic UV blockers as they are non-toxic and chemically stable under exposure to both high temperatures and UV. Inorganic UV blockers are usually certain semiconductor oxides such as TiO\textsubscript{2}, ZnO, SiO\textsubscript{2} and Al\textsubscript{2}O\textsubscript{3}. Among these semiconductor oxides, titanium dioxide (TiO\textsubscript{2}) and zinc oxide\textsuperscript{49} (ZnO) are commonly used. It was determined that nano-sized titanium dioxide and zinc oxide were more efficient at absorbing and scattering UV radiation than the
conventional size and were thus better able to block UV.\textsuperscript{46} This is due to the fact that nano-particles have a larger surface area per unit mass and volume than the conventional materials, leading to the increase of the effectiveness of blocking UV radiation. For small particles, light scattering predominates at approximately one-tenth of the wavelength of the scattered light. Rayleigh’s scattering theory states that the scattering is strongly dependent upon the wavelength, where the scattering was inversely proportional to the wavelength to the fourth power. This theory predicts that in order to scatter UV radiation between 200 and 400 nm, the optimum particle size should be between 20 and 40 nm.

12.3.8 Finishing with natural products/enzymes

Since eco-friendliness of a finishing process may determine its viability in future, researchers are increasingly looking to use natural products for finishing of textiles. One of the most versatile natural products is chitosan, which is a deacetylated derivative of chitin, which is the second most abundant polysaccharide found on earth next to cellulose. Chitin is the main component in the shells of crustaceans, such as shrimp, crab and lobster. It is also found in exoskeletons of molluscs and insects and in the cell walls of some fungi.\textsuperscript{53}

Chitin has the same backbone as cellulose, but it has an acetamide group on the C-2 position instead of a hydroxy group. The molecular structure of cellulose, chitin and chitosan are given in Figure 12.8. The presence of

![Molecular structures](image)

12.8 The molecular structures of (a) cellulose and (b) chitin and chitosan.
amino groups in chitosan imparts hydrophilicity and antimicrobial character to it. A comprehensive review of use of chitosan as antimicrobial agent mostly for cotton and other natural fibres is available.\textsuperscript{53}

Matsukawa \textit{et al.}\textsuperscript{54} used chitosan as an antistatic agent for polyester (PET) fabric. To create bonding between chitosan and PET fabric, PET was first hydrolysed in NaOH solution to generate carboxyl groups on the PET surface. The PET fabric was then treated with chitosan in 1% acetic acid by a pad-dry-cure method. Simultaneously, the chitosan films on the fabric were insolubilized by the addition of a dicarboxylic acid to cross-link the chitosan molecules. It was observed that hydrolysed PET with chitosan treatment showed a considerably lower value (6 volts) compared to those of only hydrolysed PET (2600 volts) and untreated PET (4600 volts). Also, the decreased strength of the fabric by the NaOH treatment was recovered by the chitosan treatment. A similar antistatic treatment is reported by Eom, who observed that development of electrostatic voltage decreases below one tenth of that of untreated polyester.\textsuperscript{55}

Chitosan has also been used for imparting antibacterial character to PET fibres. Jung \textit{et al.}\textsuperscript{56} made nanofibres from the blends of PET and chitosan by electrospinning onto the PET micro-nonwoven mats for biomedical applications. The PET/chitosan nanofibres were evenly deposited onto the surface, and the diameter of the nanofibres was in the range between 500 and 800 nm. The nanofibres showed significantly enhanced wetability. The antibacterial activity of the samples was evaluated utilizing the colony counting method against \textit{Staphylococcus aureus} and \textit{Klebsiella pneumoniae}. PET/chitosan nanofibre mats showed a significantly higher growth inhibition rate compared with the PET nanofibre control.

Another natural resource, neem tree has been known to have antimicrobial properties and has been used in India for many centuries. Neem (\textit{Azadirachta indica}) extract is widely used by Indian farmers to protect the cotton crop from pest and fungus. In a recent study Joshi \textit{et al.}\textsuperscript{57} have utilized this potential of neem for the development of biofunctional polyester/cotton blend fabric. Extract from the seeds of the Neem tree (\textit{Azadirachta indica}) was used for imparting antibacterial properties to the blend fabric. Durability of the neem extract on the blended fabric was ensured with the help of simultaneous application of a resin finish to the blend fabric. Resin (glyoxal/glycol) and catalyst (aluminum sulfate and tartaric acid) concentrations were optimized to get the maximum cross-linking in the fabric blend. Quantitative analysis was carried out to measure the antimicrobial activity against Gram-positive and Gram-negative bacteria. The treated fabrics inhibited the growth of Gram-positive bacteria (\textit{Bacillus subtilis}) by more than 90\% as compared with the control sample. Antimicrobial activity against Gram-positive bacteria was retained up to five machine washes and decreased thereafter. The antibacterial activity
was higher against Gram-positive bacteria as compared to Gram-negative bacteria (*Proteus vulgaris*).

Most of the synthetic fibres are non-biodegradable. This means that the used and disposed articles made from such fibres keep building up in the environment. To overcome this problem, there have been attempts to develop biodegradable polymers. However such polymers have not been able to replace conventional fibre forming polymers due to the performance and cost factors. One attractive option is to use enzymes for biodegradation of such polymers.

Enzymes are today being used at many stages in textile wet processing. Hence amylases (desizing), cellulases (biopolishing/denim processing), pectinases (cotton scouring), proteases (wool surface modification/silk degumming/laundry detergents), catalases (H₂O₂ decomposition) are in use today. Attempts have been made to degrade polyester using lipases and esterases. Although the enzyme attack on the polyester to bring about hydrolysis is a rather slow process, which excludes the possibility of enzymes being used for polyester waste disposal at least in near future, some degree of surface hydrolysis can be achieved. Hence researchers are reconciled to using enzymes for surface modification of polyester fabric,⁵⁸–⁶² which in a sense is similar to the surface hydrolysis by NaOH in that the formation of –OH and –COOH groups takes place. However the similarity ends there. As compared to alkaline treatment, the enzyme treatment takes place under much milder conditions of time, temperature and pH. The wetability obtained is higher and the effect on mechanical properties is much milder.

### 12.4 Recent advances in finishing

#### 12.4.1 Plasma processing

Textile materials are increasingly used in various industries. In these applications, the functions of textile materials are associated with phenomena such as wetting, biocompatibility, adsorption and electrical conductivity. Wetting, biocompatibility, adhesion, and many other performance properties all begin at the surface.⁶³ The properties of textile surfaces and interfaces play key roles in material processing and application technologies.⁶⁴ As with many other types of materials, the surface properties of textiles can be readily altered by the treatment of the materials with gas plasma, without impairment of their bulk mechanical properties.

The plasma treatment of textiles is a highly innovative technology in the field of superficial treatments of the material surfaces. It is based on a DBD electric discharge (dielectric barrier discharge). The use of electric
discharge creates radicals, active chemical species and ions which can modify the surfaces of textiles due to the high plasma reactivity in two ways. Surface bombardment by radicals may change the physical nature of the surface. This may result in partial destruction of scales on wool fibre surface or formation of structures or channels on other materials. The modifications applied to the surface are on the nanometre scale at room temperature, which can fundamentally change the properties like wetting, adhesion, etc. It operates under atmospheric pressure with air or inert gas and several gas mixtures. The nature of modification changes with the type of atmosphere, speed of the process, retention time, etc. The change may also be of chemical type; through formation of functional groups due the chemical reactions taking place between the radicals and the surface molecules.

In any case it generates innovative materials and new surface properties. It does not modify the bulk properties of the material. The main advantages of plasma processing over conventional chemical finishing are:

- Negligible quantity of chemicals used
- Essentially a dry process, which does not require solvents or chemicals dangerous for the environment
- The DBD technology operates at atmospheric pressure, hence no need for sealing or vacuum pumps
- Production can proceed in continuous way
- Process times are highly reduced
- They can be applied to all textile-working phases from the fibre until fabric finishing process
- Treatment on one or both fabric sides. Hence multifunctional fabrics with hydrophilic nature on one side and hydrophobic on the other can be created.

Overall, the process provides higher energy saving and a lower environmental impact.

The following functionalities can be introduced into the treated substrates:

- Increase of wetability
- Increase of hydrophilicity
- Water-repellent and oil-repellent surfaces
- Biocompatibility (antibacterial properties of adhesion and affinity to proteins and other bio molecules)
- Dyeable (increase of dyeing rate)
- Print (increase of absorbed colour quantity by the textile fibres)
- Adhesion (increase of spreading adhesion with specific compounds)
Polymer deposition in a gaseous phase as thin layer adherent to the surface (film deposition)
- Superficial ablation of the materials (etching).

12.4.2 Micro-encapsulation

Microencapsulation is a technique in which chemicals are released in a controlled manner over a long period. In this process small particles or droplets are confined in a coating to give small capsules or microcapsules. The materials inside the capsules form the core and the outside coating becomes a barrier wall. Hence the core material is isolated from hostile outside environment and can be released either slowly by diffusion through the wall or by rupture of the wall on demand. The microcapsules can then be attached to textiles, conferring various properties to them. This is an especially useful technique when the active component of a finish has high vapour pressure or when it needs to be activated only on specific occasions. The best example of a simple application of microencapsulation is carbonless copy paper. This paper has a thin coating of microcapsules on the back side. The microcapsules contain an ink which is released only under pressure. When something is written on the paper, the pressure of writing releases the ink from the layer of microcapsules and a copy of the original writing is created on a sheet kept below carbonless copy paper.

Although the industrial applications of microencapsulation began to appear almost 50 years ago in fields like pharmaceutical, cosmetics, food processing, etc., the textile industry has been a little slow in responding to this novel technique. It was only in the 1980s and 1990s that some applications started appearing on the textile front.

In the textile industry, microencapsulation has been mainly done for innovative and novel application for value addition. Much less of it is being targeted as a replacement for conventional finishing. One of the most popular applications has been addition of fragrance to the textiles by microencapsulation.

Several manufacturers have already launched such products in the market. Welbek fabrics launched its CX series of scented fabrics towards the end of the twentieth century. Similar products have also been produced by RT Dodge, Celanese International, Matsui Shikiso Chemical Co, LJ Specialities, El Dorado International Co., etc. In addition to the fragrances, another type of textiles known as Wellness Textiles based on microencapsulation are being offered. Hence there is a product: Biocap Series from Specialty Textiles Products which has microcapsules containing vitamins A, D, E and Aloe Vera. The vitamins are slowly released into the body of the wearer through the skin.
Microencapsulation is also being used for making intelligent textiles which can regulate the temperature using phase change materials (PCMs). The PCMs are compounds which have their melting temperature near the ambient temperature. When they are brought to higher temperatures, the PCMs melt and absorb heat. When they are brought down below ambient temperature, they solidify and release heat. Hence the wearer using clothes having PCMs is protected from sudden thermal shocks.

12.5 Future trends

In future one can expect some new trends in the way the textile finishing is carried out. With cost of energy, effluent treatment and competition becoming increasingly critical factors by the day, the emphasis is going to be on efficiency and cost effectiveness in finishing. With environment being a key issue, a lot of emphasis is going to be on the environmental impact of the processes.

This makes plasma processing an important technique for future textile finishing, as the process uses no water and few chemicals and causes little pollution. The role of nanotechnology in finishing will be more dominant than it is today. The main reason is that the functionalities can be created without spoiling the feel and the breathability of the garments. We are also going to witness more value addition in the products through use of techniques like micro- and nanoencapsulation.

The role of smart textiles or smart finishes will grow significantly. The finishes which can change their character according to the changes in the external stimuli, like temperature and humidity, will be more in demand. One such finish was discussed in the waterproof/breathable textiles section.

Lastly, the customers are going to look for multiple functionalities in the textile products they buy. Hence a finisher should be ready to provide easy care, soil release, antimicrobial, UV protection and breathability in a single textile product.

12.6 Sources of further information and advice

For those interested in the basic principles of textile finishing processes, there are some very good books available. The list of these is given below:


For information related to the research and latest developments in textile finishing processes, chemicals and machinery, the following journals may be referred to:

1. *Melland International* (Quarterly), Deutscher Fachverlag GmbH, Mainzer Landstr. 251, 60326 Frankfurt am Main, Germany.


4. *Journal of the Textile Institute*, The Textile Institute, 1st Floor, St James’s Buildings, 79 Oxford Street, Manchester, M1 6FQ, UK.

5. *International Dyer*, World Textile Publications Ltd, Perkin House, 1 Longlands Street, Bradford, West Yorkshire, BD1 2TP, UK.


These journals also advertise any books in the field of textiles which may be due for publication from any publisher.

### 12.7 References


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13

The impact of nanotechnology on polyesters, polyamides and other textiles

MANGALA JOSHI,
Indian Institute of Technology, New Delhi, India

13.1 Introduction

Nanotechnology is an emerging interdisciplinary area expected to have wide ranging implications in all fields of science and technology such as material science, mechanics, electronics, optics, medicine, energy, and aerospace, plastics and textiles. Although nanotechnology is still in its infancy, it is already proving to be a useful tool in improving the performance of textiles and generating worldwide interest as is evident from 1460000 hits in a Google search for the words ‘nanotechnology in textiles’. Though textile technology is about 200 years old, it can provide the bridge to making nanotechnology a commercial reality, as phenomenal efforts are being made both by the industry and academicians to use nanotechnology to create radically new textiles or to enhance the conventional textiles with greater functionality [1]. It is anticipated that the market for textiles based on nanotechnology which was approximately US$13.6 billion (2007) would rise dramatically to reach US$115 billion by 2012 [2].

13.2 What is nanotechnology?

The ‘nano’ in nanotechnology stands for one billionth of a meter, i.e. 1 nm = 10^{-9} m. ‘Nano’ is the Greek word for ‘dwarf’ and approximately 10000 times finer than a human hair. The average size of an atom is of the order of 1–2 ångstrom in radius. One nanometer comprises 10 ångstrom and hence in one nm there may be 3–5 atoms depending on the atomic radii. According to the National Nanotechnology Initiative (NNI), nanotechnology is defined as the utilization of structures with at least one nano dimension for the construction of materials, devices or systems with novel or significantly improved properties due to their nano size. In another definition of nanoscience, particles having sizes in the range of 1 to 100 nm are termed as nano particles and the application of nano-sized particles is what we call ‘nanotechnology’.
It is the desire to control properties of materials on a molecular scale that gave rise to the revolutionary field of ‘nanoscience and nanotechnology’. Nanotechnology not only produces small structures but is also an anticipated manufacturing technology, which can give thorough inexpensive control on the structure of matter. It is therefore called ‘bottom up’ technology by which bulk materials can be built precisely using tiny building blocks of nanometer size and is different from the conventional ‘top down’ technology, which starts from bulk materials and reduces to nanoscale dimension. The resultant materials therefore have fewer defects and superior quality.

The fundamental of nanotechnology lies in the fact that properties of materials change dramatically when their size is reduced to the nanometer range. For example, nanocrystalline copper is five times harder than ordinary copper with its micrometer sized crystalline structure. Since nanometer size grains, fibers and plates have dramatically increased surface area compared to their conventional size materials, the chemistry and physics of these nanosized materials is altered compared to conventional materials (Figure 13.1). Nano-elements thus manifest extremely fascinating and useful properties, which can be exploited for a variety of structural and non-structural applications.

### 13.3 Origin of nanotechnology

The concept of nanotechnology is not new as Mother Nature perfected it millions of years ago. The self-cleaning properties of lotus leaf, which are due to its carefully designed nanostructured surface and tooth and bone the natural nano biocomposites being excellent examples of nanotechnology already existing in nature.
However, the embryo of nanotechnology is believed to be the concept of ‘atomic assembly’ which was first publicly articulated in 1959 by Nobel Laureate and physicist Sir Richard Feynman, in his famous lecture titled ‘There is plenty of room at the bottom’. He stated, ‘The principles of physics so far as I can see do not speak against the possibility of maneuvering things atom by atom’. The term ‘nanotechnology’ was first coined by Norio Taniguchi at the University of Tokyo in 1974, when he defined the theoretical dimensions and tolerances in technology and also described the precision manufacturing of materials with nanometer tolerances. K. Eric Drexler, a student of MIT unknowingly appropriated the word ‘nanotechnology’ in 1986 in his famous book *Engines of Creation: The Coming Era of Nanotechnology* to describe what later became known as molecular nanotechnology (MNT). In that book, he proposed the idea of a nanoscale ‘assembler’ which would be able to build a copy of itself and of other items of arbitrary complexity. His 1991 doctoral thesis at MIT was revised and published as the book *Nanosystems Molecular Machinery Manufacturing and Computation* (1992), which received the Association of American Publishers award for Best Computer Science Book of 1992 [3].

### 13.4 Nanotechnology: applications in textiles

The textile industry is likely to be hugely impacted by nanotechnology, as research involving nanotechnology based finishes to improve performances or to create new functionality on textile materials has gained a big impetus/momentum across the globe. The US and Europe were the most active continents in the application of nanotechnology in the textile field in the beginning but more recently Asian countries account for nearly 50% of publications and patents registered in the European Patent Office, related to ‘Nano-textiles’ [4].

Incidently, the first commercial product based on nanotechnology was launched in the textile field, when Nano-Tex LLC, a subsidiary of US based Burlington Industries founded by Dr David Soane in 1998, launched its new cutting age technology ‘NanoCare’ for water repellent and stain resistant cotton fabrics, which sheds water like a duck’s feather. Later on, more and more textile companies began to invest in the development of nanotechnologies for textiles such as Swiss based Schoeller, Toray Inc from Japan, BASF from Germany, Ciba and Clariant to name a few.

The research in this area mainly centers around creating unique properties in everyday fabrics such as self-cleaning, water and oil repellency, stain proof, antibacterial, UV protective, antistatic, improved moisture regain and comfort in synthetic based textiles but all without compromising the original hand, breathability or durability of the fabric. It also shows promising applications in developing advanced textile materials such as
nanocomposite fibers, nanofibers and other nanomaterial incorporated fibers/textiles for applications in medical, defense, aerospace and other technical textile applications such as filtration, protective clothing, etc., besides a range of smart and intelligent textiles.

These developments can be broadly classified under four major areas: i.e. nanofinishing, nanocoatings, nanofibres and nanocomposites. In this chapter, the work being done under each of these domains would be reviewed under two major headings: surface modification of textiles which includes nanofinishing and nanocoating; and fiber modifications which covers nanocomposite fibers and nanofibers. Although all the nanotechnologies discussed in this chapter are with reference to textile materials both natural and synthetic, particular references to polyester and polyamide based substrates are made, wherever applicable.

13.5 Nanotechnology based surface modification of textiles

13.5.1 Nanofinishing

Nanotechnology has opened immense possibilities in the textile finishing area resulting in innovative new finishes as well as new application techniques. Particular emphasis is on making chemical finishing more controllable and durable with significantly enhanced functionality by incorporating various nanoparticles or creating nanostructured surfaces, which led to an unprecedented level of textile performances such as stain resistant, hydrophilic, antistatic, wrinkle resistant and shrink proof abilities [5–7].

Nanofinishes are generally applied in nanoemulsion form, which enables a more thorough, even and precise application on textile surfaces. They are generally emulsified into either nanomicelles, made into nanosols or wrapped in nanocapsules that can adhere to textile substrates easily and more uniformly. Ideally nanoparticles or the molecules of these nanofinishes can be brought individually to designated sites on textile surfaces in a specific orientation and trajectory through thermodynamic, electrostatic or other approaches. Since nanoparticles have a large surface area to volume ratio and high surface energy, they have better affinity for fabrics. Therefore these finishes are more durable, effective and do not adversely affect the original handle and breathability of the fabric. A range of different textile products and finishes based on nanotechnology has already been launched in the market and these are summarized in Table 13.1. The recent developments in nanofinishing on textiles are reviewed and briefly described here.
<table>
<thead>
<tr>
<th>S. No.</th>
<th>Name of the finish</th>
<th>Company</th>
<th>Properties</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Resists Spills™</td>
<td>Nano-Tex</td>
<td>Water-repellent and stain resistant</td>
<td><a href="http://www.nanotex.com">www.nanotex.com</a></td>
</tr>
<tr>
<td>2.</td>
<td>Coolest comfort™</td>
<td>Nano-Tex</td>
<td>Impart superior wicking properties to a previously hydrophobic synthetic or resin treated cotton</td>
<td><a href="http://www.nanotex.com">www.nanotex.com</a></td>
</tr>
<tr>
<td>3.</td>
<td>NanoCare®</td>
<td>Nano-Tex</td>
<td>Helps stains to wash out easily</td>
<td><a href="http://www.nanotex.com">www.nanotex.com</a></td>
</tr>
<tr>
<td>6.</td>
<td>NanoSphere®</td>
<td>Schoeller Textil AG</td>
<td>Repels water and oil drops and prevents soil particles from attaching themselves on the fabric surface</td>
<td><a href="http://www.schoeller-textiles.com">www.schoeller-textiles.com</a></td>
</tr>
<tr>
<td>8.</td>
<td>Nuva FW</td>
<td>Clariant</td>
<td>Excellent oil and water-repellent effects on textiles made from synthetic and cellulosic fibres</td>
<td><a href="http://www.textiles.clariant.com">www.textiles.clariant.com</a></td>
</tr>
</tbody>
</table>

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13.5.2 Water and oil repellent (hydrophobic) nanofinishes

A number of water and oil repellent finishing chemicals are available commercially which could be classified into wash resistant or not wash resistant on one hand and fluorocarbon containing or non-fluorocarbon based finishes on the other hand. Silicon based water proofing is also used either as such or in combination with fluorocarbon based agents. However the current market demands are for non-fluorocarbon based finishes because of growing environmental consciousness and increasingly strict legal regulations. Besides this, the durability of finish and retaining the original feel and strength of the fabric are also important criteria. Nanotechnology based novel and innovative nanofinishes seem to be closer to achieving these consumer demands.

The premier range of NanoCare® and NanoPel® nanofinishes marketed by NanoTex Inc. USA are the next generation easy care finishes based on nanotechnology. These finishes which come under the Resist Spills™ Category protect the fabric against both water and oil based liquid stains/soils [8]. Tiny whiskers aligned by proprietary ‘spines’ are designed to repel liquids and are attached to the fibers utilizing molecular ‘hooks’. These whiskers and hooks are minute, in fact no more than 1/1000th the size of a cotton fiber. These whiskers cause the liquids or semisolids to roll off the fabric thus causing minimal staining and can be removed with simple washing (Figure 13.2). Since the attached whiskers are of nanoscale

![NanoCare Technology: Tiny whiskers attached to individual cotton fibers cause the liquid to roll off.](image)
size, they do not affect the hand, breathability of fabric and can withstand 50 home launderings. The Resist Spills™ protection is applied to fabrics at mill level during finishing stage and requires no special machinery or processing conditions. The treatment can be applied either by padding or dip or dry coating. This nanofinish can be applied to natural fibers such as cotton, wool and silk as well as synthetics such as polyamides, polyesters, acrylic, etc. Leading garment-manufacturing brands such as Burlington, Galey & Lord, Dan River, Eddie Bauer and Lee have used this licensed technology for their NanoCare® range of commercial products. The large-scale production of shirts, shirting fabrics has also been licensed to several South Asian countries including India.

What are ‘whiskers’?

The original research demonstrated and produced by Dr David Soane forms the basis of ‘NanoCare’ range of finishes [7, 9–11]. The tiny whiskers ‘10–100 nm’ in dimension are nothing but low molecular weight nanosized multifunctional molecules, which are capable of forming covalent and non-covalent bonds with cellulosic and protein fibers. These molecules are either block or graft copolymers, having multiple functional groups such as binding groups, hydrophobic groups hydrophilic and oleophilic groups. These nanosized molecules react with reactive groups such as hydroxyl, carboxyl or amino present in the surface of the fabric without blocking its pores. These tiny whiskers modify the surface properties of the treated fabric and improve water repellency, grease repellency, soil resistance and detergent free washing without affecting its strength, abrasion resistance and breathability. Due to the multiplicity of bonds and the ability of the molecules to easily diffuse into the fiber because of small nanosize, the durability of the finish is much better than the conventional fluorocarbon acrylate polymer based finish (Figure 13.3).

13.5.3 Superhydrophobic: self-cleaning nanofinishes

Many plants in nature including the lotus leaf exhibit unusual wetting characteristic of superhydrophobicity. A super hydrophobic surface is the one that can bead off water droplets completely; such surfaces exhibit water droplet advancing angles of 150 degrees or higher (Figure 13.4). In addition their contact angle hysteresis is very low (receding contact angle only a few degrees less than the advancing contact angle), producing a surface on which water droplets just roll off. A self-cleaning surface thus results since the rolling water droplets across the surface can easily pick up the dirt particles to leave behind a clean surface.
This fascinating self-cleaning property of lotus leaf has been accomplished by nature through the use of a surface roughness in the combination with a highly hydrophobic surface. The surface topography of lotus leaf presents two different scales to the outside environment and is textured with 3–10 micron sized hills and valleys that are decorated with nanometer sized particles of a hydrophobic wax like material (Figure 13.5). The hills and valleys ensure that surface contact area available to water is very low while the hydrophobic nanoparticles prevent water into the valleys. The net result is that water cannot wet the surface and therefore forms nearly spherical water droplets, leading to superhydrophobic surfaces [12]. The increasing surface roughness due to multiplicity of microscale or nanoscale projection or cavities provides a large geometric area for a relatively small projected area. Cassie and Baxter [13] were the first to explain that water repellency of rough surfaces was due to the air enclosed between the gaps in the surface, which minimizes the substrate to water interface. Hence water does not spread and forms a droplet, which easily rolls off.

Taking the inspiration from nature there have been several approaches researched to create superhydrophobic surfaces on textiles, which mimic
the nanostructured lotus leaf and therefore exhibit self-cleaning properties. Nano Sphere®, a lotus effect based textile finish has been developed, patented and commercialized by Schoeller Texil AG of Switzerland [14]. This water and oil repellent finish is based on a ‘guest–host’ system and uses a dispersion system comprised of the dispersed phase ‘guest’ and the dispersant ‘host’ components [15]. In the final finishing layer, self-organization of the ‘guest’ and the ‘host’ components leads to the ‘guest’ component concentrating at the upper surface of the finish layer and forming columnar structures similar to the microrough surface exerting the ‘lotus effect’. The partial phase separation of the ‘guest’ and ‘host’ system results in the accumulation of the hydrophobic or oleophobic ‘guest’ components at the surfaces. The phase separation between the finishing layer and the surrounding air results in the dramatically enhanced water and oil repellent property of the fabric. The dispersant is an emulsion of a gel-forming component (polymeric binder) and a water and/or oil repellent components. The third component, which is the dispersed phase, is comprised of either micro sized waxes (0.1–50 micron) or hydrophobic silica particles (5–50 nm). The finish can be applied on textile material by conventional industrial application techniques such as padding, coating, spraying or foaming and can be applied to both natural, i.e. cotton, viscose, polyester/cotton blends as well as wholly synthetic polyester or polyamide textile substrates.
Superhydrophobic silica coating film on cotton substrates, which are transparent and durable have been reported by W. A. Daoud and coworkers of the Hong Kong Polytechnic University using low temperature sol-gel coating based on a low temperature process [16]. The coatings were produced via cohydrolysis and polycondensation of a hexa decyl trimethoxy silane (HDTMS), tetra ethoxy silicate (TEOS) and 3-glycidyloxy propyl trimethoxy silane (GPTMS) mixture. The superhydrophobic nanocomposite coating enhances the contact angle of cotton to 140°, whereas before coating cotton has 0° contact angle as it is superhydrophilic in nature. This nanocomposite coating has new applications in daily use material such as plastics or textiles and is an ecofriendly substitute for fluorocarbon based water repellant finish. There is less than 5% decrease in textile strength and tearing strength. The air permeability of the fabric remains unchanged. The washing durability of the coatings is also good which is attributed to linking ability of GPTMS that promotes a high level of adhesion at the interface.

Yu et al. [17] recently reported another superhydrophobic complex coating for cotton fabrics based on silica nanoparticles and perfluorinated quaternary ammonium silane coupling agent (PFSC), using a sol-gel process. Silica nanoparticles in the coating made the textile surface much rougher and PFSC, on the top layer of the surface, lowered the surface free energy. Textiles coated with this coating showed excellent water repellant property and water contact angle increased up to 145° from 133° for cotton fabrics treated with pure PFSC, without silica sol pretreatment. The silica sol and PFSC were applied to cotton fabrics by conventional pad-dry-cure process. Superhydrophobic coatings have also been reported with many other hydrophobic film-forming nanoparticle compositions such as Al₂O₃ gel, ZnO nanoparticles, aligned carbon nanotubes, CaCO₃ loaded hydrogel spheres and POSS nanoparticles as reported in an excellent review on nanofinishing by Gulrajani [7].

However durability against laundering and abrasion is important for textile applications. A combination of chemical and mechanical treatments can be used to create super hydrophobic nanostructures on the surface of textile materials. Mechanically roughened surfaces became an integral part of the textiles and are expected to be more durable. Mechanical treatments such as calendering, embossing, etching, schreinering, sueding, sanding or emerizing have also been used to create mechanically abraded micro roughened surfaces on textiles [7].

13.5.4 Photocatalytic self-cleaning nanofinishes

Dr John Xin and Dr Walid Daoud of the Hong Kong Polytechnic University’s Nanotechnology Centre for Functional and Intelligent Textiles and
Apparel [18–20] developed a process for the sol gel coating on textile substrates at low temperature. They also claimed that photocatalytic self-cleaning properties could be imparted to the coated fabric on coating cotton with TiO$_2$ nanoparticles that are about 20 nm in size (Figure 13.6). The coating solution comprises a translucent sol prepared with a TiO$_2$ precursor (TTIP, titanium tetraisopropoxide), ethanol and acetic acid in a molar ratio of 1:100:0.5 respectively. The principal chemical reactions are summarized as follows:

$$\text{Ti(OPr)}_4 + 4 \text{EtOH} \rightarrow \text{Ti(OEt)}_4 + 4 \text{Pr OH}$$

$$\text{Ti(OPr)}_4 \text{ or Ti(OEt)}_4 + \text{H}_2\text{O} \rightarrow \text{Ti(OH)}_4 + 4\text{PrOH or 4EtOH}$$

$$\text{Ti(OH)}_4 \rightarrow \text{TiO}_2 + 2\text{H}_2\text{O}$$

The fabrics are coated with this sol using conventional pad dry cure process. The coated fabrics show discoloration of red wine and coffee stains on irradiation of samples in a Sun test solar simulator. The photocatalytic activity of semiconductor oxides such as TiO$_2$ is attributed to promotion of an electron from the valance band (VB) to the conduction band (CB) brought about by the absorption of a photon of ultra band gap (~3.2 eV) light whose energy ($h\nu$) > EBG. EBG is the energy difference
between electrons in the VB and CB. In the presence of oxygen and/or water, superoxide (O\textsuperscript{2−}) and/or hydroxyl (OH\textsuperscript{•}) radicals are formed, which attack adsorbed organic species on the surfaces of TiO\textsubscript{2} and decompose them. The mechanism suggested by the investigators [21] for this discoloration is shown in (Figure 13.7a, b).

The decomposition of stain, which is an organic compound, goes through a cation intermediate (stain+) leading ultimately to the production of CO\textsubscript{2}. The electron generated in the process is injected into TiO\textsubscript{2} and the conduction band starts the oxidative radical chain, leading to discoloration of the...
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The nanotitania coated fabrics maintain their antibacterial property up to 55 washes/home launderings and UV protection characteristics up to 22 washes. This has been attributed to the formation of interfacial bonding through a dehydration reaction between the cellulosic hydroxyl groups of cotton and the hydroxyl group of titania [22].

13.5.5 Hydrophilic nanofinishes

The poor moisture absorption property of synthetic fabrics such as polyester and polyamides limits its applications in the apparel sector. The new range of hydrophilic nanofinishes Cotton Touch™ and Coolest Comfort™ commercialized by NanoTex, USA makes the synthetic fabric look and feel like cotton [8].

*Cotton Touch™*

This is a naturally soft fabric enhancement designed to make synthetic fabric look and feel like cotton. It is available as Nanotouch® and gives a durable cellulose wrapping over synthetic fibers such as polyester and polyamides. Cellulosic sheath and synthetic core together form a concentric structure to bring overall solutions to the drawbacks of synthetics such as static discharge, harsh handle and glaring luster. It can also last 50 launderings and is expected to eliminate the decline in demand of synthetic microfiber and broaden the use of synthetics to new applications.

*Coolest Comfort™*

This fabric provides breakthrough moisture wicking to draw moisture away from the body while drying quickly. The popular product in this category is Nano Dry®. It improves the moisture absorption of polyamides and polyesters making them hydrophilic and comfortable. The main applications are in sportswear and close-to-body garments that require perspiration absorbency. The finish lasts 50 launderings.

13.5.6 Antibacterial nanofinishes

Silver (Ag) or silver ions (Ag⁺) have long been known to have strong inhibitory and bactericidal effects as well as a broad spectrum of antimicrobial activities [23]. The inhibitory effects of silver ion/metal on bacteria have been attributed to the interaction of silver ion with thiol group in bacteria as well as to the oxidative destruction of microorganisms in aqueous medium [24–25] (Figure 13.8).
Even though metallic silver has antimicrobial properties, it is expected that the conversion of bulk silver to nanoparticles, i.e. ‘nano Ag’ will have very high specific area that may lead to high antimicrobial activity compared to bulk Ag metal. A range of antimicrobial textile finishes and products have been reported and quite a few have been commercialized, which are based on superior antimicrobial properties of silver in nanoform and are summarized in Table 13.2. Several methods have been devised to synthesize silver nanoparticles and reviewed [7]. These include chemical reduction, chemical/photo reduction in reverse micelles, or in microemulsions, radiation induced chemical reduction and mechano-chemical reaction between AgCl and Na in a ball mill. In a chemical method of producing highly concentrated stable dispersion of nano sized silver particles, silver nitrate is reduced with ascorbic acid to precipitate silver metal in acidic solution as per the following reaction [26]:

$$2\text{Ag}^+ + \text{C}_6\text{H}_8\text{O}_6 \rightleftharpoons 2\text{Ag} + \text{C}_6\text{H}_6\text{O}_6 + 2\text{H}^+$$

Silver ions and nanoparticles both have to be stabilized, as they are prone to discoloration on exposure to light. The stabilization of silver nanoparticles is achieved either by coating, encapsulation or complex formation between the lone pair of electrons on N and Ag atoms. All amines having a free lone pair of electrons can stabilize Ag ions/nanoparticles. Polyvinyl pyrrolidone (PVP) is one such example [27].

Antimicrobial yarns made from cotton, linen, silk, wool, polyester, nylon or their blends have been produced by immersing them in silver nanopar-
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ticle containing solutions. The treated yarns are yellow in color and contain 0.4–0.9% by weight of silver. The treated yarns show effective antimicrobial activity against various bacteria, fungi, etc., and the activity remains even after 100 washes with neutral soap. Nano silver particles containing antimicrobial dressings have been incorporated in wound care and have gained wide acceptance in medical industry, as a safe and effective means of controlling microbial growth in the wound, often resulting in improved healing.

13.5.7 UV protective nanofinishes

Semiconductor oxides such as TiO$_2$, ZnO, SiO$_2$ and Al$_2$O$_3$ are known to have UV blocking property [29–30]. It is also known that nanosized TiO$_2$ and ZnO particles are more efficient at absorbing and scattering UV radiation than the conventional size particles and thus were better able to block UV radiation as they have a much larger surface area to volume ratio and also the fact that Rayleigh Scattering due to nanoparticles is much reduced as it is inversely proportional to the fourth power of their sizes. This theory predicts that in order to scatter UV radiation between 200 and 400 nm, the optimum particle size should be between 20 and 40 nm [29]. Moreover inorganic UV blockers are advantageous as compared to organic UV absorbers, as they are less toxic and chemically stable under exposure to high temperature and radiation.

Much effort has been made on the application of UV bulking treatment to fabrics using nanotechnology [31–35]. UV blocking treatments for cotton fabric have been developed using the sol-gel method by Xin and coworkers [34]. A thin layer of TiO$_2$ nanoparticle is formed, on the surface of treated cotton fabric, which provides excellent UV protection, the finish is durable up to 50 home launderings. Apart from TiO$_2$, ZnO nanorods of 10 to 50 nm in length were also applied to cotton fabric to provide UV protection. The rods exhibited excellent UV protection [36].

13.5.8 Antistatic nanofinishes

Synthetic fibers such as nylon and polyester are prone to static charge accumulation as they absorb less water. It has been reported that nanosized TiO$_2$, ZnO whiskers, nanoantimony-doped tin oxide (ATO) and silane nanosol could impart antistatic properties to synthetic fibers [37–39]. ZnO and TiO$_2$ nanoparticles are electrically conductive materials and help dissipate the static charge in these fibers. Silane nanosol improves antistatic properties as the silane gel nanoparticles absorb moisture due to $-\text{NH}_2$ and $-\text{OH}$ groups present. W. L. Gore and Associates GmbH have combined
<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Company</th>
<th>Product name</th>
<th>Type of product</th>
<th>References</th>
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<tr>
<td>1.</td>
<td>NanoHorizons</td>
<td>SmartSilver™</td>
<td>Additive</td>
<td><a href="http://www.nanotech-now.com">www.nanotech-now.com</a></td>
</tr>
<tr>
<td>2.</td>
<td>ABCNanotech</td>
<td>SARPU</td>
<td>Finishing/coating solution (nanosized silver dispersed in liquor)</td>
<td><a href="http://www.abcnanotech.com/">http://www.abcnanotech.com/</a></td>
</tr>
<tr>
<td>5.</td>
<td>JR Nanotech</td>
<td>SoleFreshT</td>
<td>Dispersion containing 0.3%w/w nanosilver</td>
<td><a href="http://www.jrnanotech.com">http://www.jrnanotech.com</a></td>
</tr>
<tr>
<td>7.</td>
<td>NanoGap</td>
<td>Silver nanoparticles</td>
<td>Powders or in organic or aqueous solutions</td>
<td><a href="http://www.nanogap.es">http://www.nanogap.es</a></td>
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</table>
nanotechnology and PTFE (Dupont’s Teflon) to develop an antistatic membrane for protective clothing. Gore-Tex® based works wear protects the wearer from electrostatic discharges [40]. Electrically conductive nanoparticles are durably anchored in the fibers of the membrane, creating an electrically conducting network that prevents the accumulation of static charge.

13.6 Nanocoatings

Nanostructured surfaces are of great interest, due to their large surface area, which might yield high functionality. Nanocoating refers to the covering of materials with a layer on the nanometer scale (10–100 nm in thickness) or covering of a nanoscale entity to form nanocomposite and structured materials. There are numerous coating procedures that are widely used including chemical vapor deposition, plasma assisted techniques, self-assembly, layer by layer coating, dip coating, sol-gel coating and electrochemical deposition mostly on planar substrates, i.e. metal, ceramic or polymer. Coating in solution is generally performed using either precursor molecules or preformed nanoparticles to form the layer. Electrostatic interaction, hydrogen bonding and covalent bondings are some of the associated forces between the coating and the material being coated. Nanocoatings on textiles have recently been explored using mainly processes such as plasma-assisted polymerization, self-assembly, sol-gel nanocoating and electrochemical deposition which will be briefly reviewed in this section.

13.6.1 Self-assembly

Toray Industries, Inc. have succeeded in developing a ‘nano-scale processing technology’ that allows the formation of molecular arrangement and molecular assembly necessary to bring out further advanced functionalities in textile processing [41]. This ‘nano-scale processing technology’ named ‘NanoMATRIX’ forms the functional material coating (10–30 nm) consisting of nano-scale molecular assembly on each of the monofilaments that forms the fabric (woven/knitted fabric) (Figure 13.9). NanoMATRIX is based on the concept of ‘self-organization’ by controlling the conditions like temperature, pressure, magnetic field, electrical field, humidity, additives, etc. It is possible to control the state of molecular arrangement and/or assembly of functional materials on each of the monofilaments in nanoscale sizes precisely by controlling the interaction and responses between the functional material to be coated and the fiber material (polymer). The application of this technology is expected to lead to the development of new functionalities as well as remarkable improvements in the existing
functions (quality, durability, feel, etc.) without losing the fabric’s texture.

13.6.2 Plasma assisted nanocoatings

Plasma polymerization enables deposition of very thin nanostructured coatings (<100 nm) via gas phase activation and plasma substrate interactions. This dry and ecofriendly technology offers an attractive alternative to replace wet chemical process steps for surface modification (finishing) of textiles. Plasma polymerization can impart a wide range of functionalities such as water repellency, hydrophilicity, dyeability, conductivity and biocompatibility due to the nanoscaled surface modification of textiles and fibers. The advantages over conventional wet chemical processing are that it needs a very low material and low energy input, hence is environmentally
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friendly, it does not affect the bulk properties of textiles and fibers such as feel (touch), handle, optical properties and mechanical strength. Moreover, these plasma-assisted coatings are more durable as compared to other surface modification techniques such as wet processes, radiation or simple plasma activation because nanoscaled plasma polymer coatings get covalently attached or bonded to textile surfaces [42, 43].

What is plasma polymerization?

Plasma is considered the fourth state of matter and consists of a wide variety of active species such as electrons, ions, radicals, excited species, vacuum and ultraviolet radiation. Surface treatment of textiles is usually carried out by non-equilibrium plasmas, which are excited by the electric field. The excited electrons in the plasma have the appropriate energy to excite, dissociate and ionize atoms and molecules. Monomer gases when introduced into the plasma reactor, enter an active zone where excitation and dissociation process are taking place to generate reactive species and then travel to a passive zone yielding recombination and stable products which get deposited on substrates, electrode or wall as thin plasma polymerized coatings (Figure 13.10). Thus plasma polymerization is a radical
dominated chemical vapour deposition (plasma CVD) process, thought to result in macromolecule formation i.e. mainly amorphous, and more or less cross-linked structures.

Plasma polymerization is performed using different kinds of plasma polymerizable gases (monomers) such as hydrocarbons (i.e. methane, ethylene or acetylene) or organosilicon monomers (i.e. silicone (SiH₄) or silicon tetrachloride (SiCl₄)). There is a difference between conventional and plasma polymerization, as some gases (like methane) which can be plasma polymerized do not undergo polymerization by the conventional route. A plasma polymer typically results from a rivaling etching and deposition process depending on the plasma species present during film growth yielding a more or less cross-linked structure [44–46].

Plasma equipment

There are two types of plasma, i.e. high temperature plasma and low temperature plasma. High temperature plasma is found at atmospheric pressure in either its man made form such as plasma torch or its natural form such as lightning. Low temperature (also called low pressure) plasma techniques are mostly suitable for surface reaction, cleaning, etching and plasma polymerization [46].

Most of the earlier work on plasma processing of textiles is based on low-pressure plasmas operating between 0.1 Pa and 100 Pa, which can be activated by direct current (DC), alternative current (AC) and radio frequency or microwave (MW). However, high-pressure atmospheric plasmas, such as corona and dielectric barrier discharge (DBD), are of special interest to textile industries because of easier processibility. However, it is difficult to obtain controlled deposition conditions or uniform coating through high-pressure atmospheric plasma. This is because the plasma can be ignited only within smaller volumes, hence non-coated parts are also taking part in external interaction. Moreover, the atmospheric plasma is mostly filamentary in nature, thus leading to moving hot spots. Research is ongoing to overcome these limitations [47, 48, 49].

M/s EMPA, a Swiss based company specializing in this area, have developed a low-pressure plasma reactor for plasma-polymerized coatings. At low pressure (1–100 pa) the plasma zone is well defined and the mean free path lengths are high enough to allow the penetration of textile structure by energetic particles as well as by long living radicals, thus enabling textile fibers up to several fiber layers in depth to interact with external plasma media. EMPA has developed both a semi-continuous web coater (up to 63 cm in width) and a continuous fiber coater besides several batch reactors. The reactors have been typically designed for textile widths up to 120 cm but larger widths of up to 4 m are also possible. The textiles are
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led through plasma regions and adapted to the required process, which comprises usually a cleaning step followed by a plasma deposition step. As compared to plasma polymerization on textile fibers or web, it is easier to do it on fibers as it needs smaller reactors and the process can be more continuous. Prior to plasma polymerization, the substrate needs to be cleaned as well as activated for better adhesion with the coating [46, 48].

Plasma polymerized functional nanocoatings

The earliest plasma treatments on textiles date back to the 1960s and focused on improving the wettability of fabrics through plasma activation. Plasma treatment in presence of non-polymerizable gases such as N₂, NH₃, O₂, H₂O, CO₂ or others generates oxygen or nitrogen-containing functional groups such as carboxyl (–COOH), amino (–NH₂), hydroxyl (–OH), or ethylene oxide (EO) units (–CH₂–CH₂–O–) on synthetic fabrics such as (PP, PET, cellulose acetate, and others) [48, 49]. The positive effects of this process are cleaning, etching, cross-linking, formation of radical sites and formation of functional groups which are not only able to enhance the binding sites in subsequent process steps of coating, dyeing, lamination and fiber reinforced composites but also to increase the surface energies of textile surfaces so as to make them more hydrophilic [41, 46, 50]. However this type of plasma activation is prone to aging due to internal reorientation effect as well as external influences. Aging can however be minimized by deposition of functional plasma polymers. Thus the internal reorientation process gets hindered by the cross-linked film network, which can be achieved by mixing hydrocarbon polymerizable gases with non-polymerizable gases, that is oxygen and nitrogen, etc. Rivaling process of etching and plasma polymerization yield highly functionalized and permanent plasma polymer coatings on textile substrates. Plasma polymerization can be performed with any kind of hydrocarbon monomer (in gaseous form), mainly acetylene, ethylene or methane are used on textile substrates. The achievable fiber properties depend on degree of cross-linking in the otherwise amorphous polymer coating deposited. The deposition of plasma polymerized coatings with well-defined penetration properties can be achieved by controlling the energy input into the active plasma zone which can be of use in controlled drug release from textile substrates. The coated surfaces also possess hydrophobic property [46].

Plasma polymerization of ethylene monomer on PET fibers was found to increase the interfacial adhesion strength between PET fibers and PE matrix in fiber reinforced composites [51]. Modification of polyester cords by low temperature plasma at atmospheric pressure in the presence of
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Nitrogen and butadiene resulted in homogeneously coated cord surfaces. The plasma polymerized cords showed values of static and dynamic adhesion comparable to chemical treatment based on resorcinol formaldehyde latex (RFL) [52].

Plasma polymerization of organosilicon compounds is reported to impart good dielectric property, thermal stability, scratch resistance, lowered friction, flame retardant and barrier properties as well as adjusting the wetability of textiles [46, 53, 51]. Hexamethylene disiloxane \( (\text{HMDSO}, (\text{CH}_3)_3\text{Si-O-Si(CH}_3)_3) \) is the best common monomer, it is nontoxic and enables the deposition of siloxane coatings at low temperature. HMDSO derived plasma polymers are used for the hydrophobization of cotton, showing water contact angles up to 130° without influencing the water vapour transmission through the fabric [54]. Low-pressure plasma polymerization of unsaturated fluoro hydrocarbons, i.e. \( \text{C}_2\text{F}_6, \text{C}_4\text{F}_8 \) on selected textiles has been industrially performed using a semi-continuous process to impart stain repellent properties on fabrics. Oil repellency grades of 4–5 were achievable in short treatment times (30–60 sec), which is superior to commercially available Scotch Guard finished samples. The softness, feel, color, permeability, abrasion resistance, water performance and friction coefficient properties of original fabric were unaltered by these nanoscaled ultra-thin (<100 nm) plasma coatings [55].

Multifunctional acrylic-like coatings on fibers such as polyester or polyamide have been obtained by low-pressure plasma polymerization using acrylic acid as a monomer. Acrylic coatings were found to improve their wettability, dyeability (using acid dyes) and soil resistance [56] and also improve the cell adhesion for tissue engineering [57]. The wrinkle recovery angles also improved by about 13% and no undesirable decrease in tensile strength was observed [58].

Ceramic Nanocoatings such as \( \text{TiO}_2 \), which are known for their superior hydrophilicity and photocatalytic activity, have also been obtained using plasma polymerization in presence of tetra isopropyl orthotitanate (TTIP) and oxygen as gases at low temperature (<100°C) [59]. The incorporation of nanoparticles into wet chemical coatings presents some difficulties such as agglomeration or non-uniform distribution of nanoparticles at surfaces. An in situ plasma polymerization/co-sputtering process has been used to embedded nanoparticles with homogeneous size and spatial distribution. Hegemann and coworkers [60] have recently used low pressure RF plasma discharges using acetylene mixed with ammonia to obtain interconnected nanoporous highly crosslinked coatings on textile fabrics. Co-sputtering of a silver target with Ar enabled the in situ incorporation by Ag nanoparticles within the functional plasma polymerized coating. A homogeneous distribution of Ag nanoparticles present at the coating
surface was obtained and imparted antimicrobial properties to the coated substrate (Figure 13.11). A range of other functionality can be introduced onto fabrics by depositing uniform mono-dispersed nanostructured particles such as TiO$_2$ for catalytic and ferrous oxide for magnetic properties on the surface of fibers using this plasma polymerization technique developed by M/s EMPA.

Thus plasma polymerization can be applied to all forms of textile products such as fabrics and fibers. The characterization of the plasma polymerized nano-coatings is mainly affected by the type of precursor used, precursor flow rate, and plasma generator power applied to reactor, generator frequency and the exposure time of the substrate to plasma conditions. The up-scaling of plasma technology to industrial scale for textile applications is the major challenge faced by the researchers and technologists. Low-pressure plasma processes are still the state of the art technology, as effects produced by atmospheric plasma are comparatively weak and non-uniform. The other issues of concern are the efficiency of plasma polymerization process in terms of deposition rates and the right process speeds, so that they can be integrated with the current textile pro-
duction lines. High investment cost and requirement of vacuum technology further limits the present application of this technology at an industrial scale to only niche textile products [46].

13.6.3 Sol-gel nanocoatings

The sol-gel process is a widely used technique to coat surfaces with nanoscale entities, i.e.: nanoparticles and finds application in a variety of areas ranging from catalysis, electronics, biomedical engineering and material science [61]. The sol-gel process involves inorganic precursors (a metal salt of an organo metallic molecule) that undergoes various reactions resulting in formation of a three dimensional molecular network. A common example is the hydrolysis and condensation reaction of metal alkoxides to form larger metal oxide molecules:

\[
\begin{align*}
\text{Hydrolysis} \\
&M(OR)_4 + H_2O \rightarrow HO-M(OR)_3 + ROH \rightarrow M(OH)_4 + 4ROH \\
\text{Condensation} \\
&(OR)_3M-OH + HO-M(OR)_3 \rightarrow (OR)_3M-O-M(OR)_3 + H_2O \\
&(OR)_3M-OH + RO-M(OR)_3 \rightarrow (OR)_3M-O-M(OR)_3 + ROH
\end{align*}
\]

where M represents the metal and R represents alkyl group.

Low temperature sol-gel based nanocoatings have been recently used to deposit layers of metal oxide nanoparticles on textile fabric surfaces to impact specific functionality, i.e.: nanotitania (TiO\textsubscript{2}) for photocatalytic activity, nanosilica (SiO\textsubscript{2}) for superhydrophobic lotus leaf effect [7, 16, 17, 18–20] as also discussed in section 13.5.1 on ‘Nanofinishing’.

One of the major challenges in sol-gel nanocoatings is controlling the adhesion of the inorganic coatings to the substrates and also the uniformity of coating thickness. Hence reaction conditions need to be carefully monitored and controlled to allow reproducible coatings.

13.6.4 Layer-by-layer nanocoatings

Layer-by-layer (l-b-l) nanocoating technique can be utilized to fabricate thin film coatings, with molecular level control over film thickness and chemistry. A coating of this type can be applied to any surface amenable to the water based l-b-l adsorption process used to construct polyelectrolyte multi-layers including inside surfaces of the complex objects [62]. L-b-l assembly process involves sequential adsorption of oppositely charged polyelectrolytes on a solid support that results in multi-layer
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Apparatus is required for the l-b-l process and nanocoatings can be prepared under mild physicochemical conditions. L-b-l technique has been used to create superhydrophobic surfaces which can be applicable as anti-fouling, self cleaning and water resistant coatings for micro-fluids channels and bio sensors to name a few [62, 63].

20–50 nm thick polymer surface layers, which increased the interaction antimicrobial nylon and silk fabrics by l-b-l deposition of silver nanoparticles. The heated fabrics are sequentially dipped in a solution of PDADMAC [poly (diallyl dimethyl ammonium chloride)] and silver nanoparticles, activity [65].

Because of a unique cross-section as well as chemical heterogeneity of its surface. Cationic cotton surface has been coated with alternate layers of polyelectrolytes, i.e. poly (sodium 4-styrene sulphonate) and poly (allyl-amine hydrochloride) using l-b-l technique [66]. Our research group at

Nanocoatings on fibers or textiles is to date a relatively unexplored area with very few cited reports [64–66]. Systematic modification of the surface of lignocellulosic fibers was performed by L-b-l nanocoating process to produce negative and positively charged fibers. The fibers are coated with 20–50 nm thick polymer surface layers, which increased the interaction between the fibers during paper formation and helped to obtain stronger paper from virgin fibers [64]. Dukas and coworkers have reported antimicrobial nylon and silk fabrics by l-b-l deposition of silver nanoparticles. The heated fabrics are sequentially dipped in a solution of PDADMAC [poly (diallyl dimethyl ammonium chloride)] and silver nanoparticles, which formed a colored film of nanoparticles having strong antimicrobial activity [65].

Cotton fibers offer unique challenges to the deposition of nanolayers because of a unique cross-section as well as chemical heterogeneity of its surface. Cationic cotton surface has been coated with alternate layers of polyelectrolytes, i.e. poly (sodium 4-styrene sulphonate) and poly (allyl-amine hydrochloride) using l-b-l technique [66]. Our research group at

13.12 Schematic of layer-by-layer nanocoatings through self-assembly [65].
IIT Delhi has recently investigated this technique to deposit nanocoatings on textile surfaces using polyanion (PAH) and polycation chitosan on cotton fabric [67].

### 13.7 Nanocomposite coatings

A glance at the literature available shows interesting applications of polymer nanocomposites as coatings with attractive combinations of properties not achievable by neat polymeric conventional coatings. Both scientists as well as industry are actively engaged in exploring the potential applications of polymer nanocomposites as advanced coating options and some of the developments in this area are described here.

Organic-inorganic hybrid nanocomposite coatings frequently exhibit unexpected hybrid properties synergistically derived from the two chemically different components. Coatings and films based on these organic-inorganic hybrid materials are endowed with exceptional mechanical and thermal stability, allowing small thickness and making them very interesting for various applications. The organic component in the matrix offers the advantages of mechanical toughness and flexibility while the inorganic component, i.e. nanoclays or metal/metal oxide nanoparticles provides hardness, thermal stability and gas barrier properties. It has been reported that by incorporation of nanoparticles, the thin film coatings have stronger bonds and better flexibility with little cost difference. These coatings are smoother, stronger and more durable. When used on products, the results range from scratch resistant and self-cleaning surfaces to moisture absorbing clothing [68, 69]. Many companies around the world are using the properties of nanoparticles and are incorporating them within their coatings.

Novel polyurethane/MMT (clay) based nanocomposites as coatings for inflatables have been explored in an ongoing research project at the Department of Textile Technology, the Indian Institute of Technology, Delhi by Joshi *et al.* [70]. The coated fabrics showed improved gas barrier property without affecting the transparency and tear strength. Clays are believed to increase the barrier properties by creating a tortuous path that retards the progress of gas molecules, i.e. gas diffusion through the matrix resin as reported [71]. Recently Wilson Sporting Goods have introduced the Wilson Double Core Tennis Ball. The manufacturers claim that these balls retain their pressure and bounce for twice as long as conventional tennis balls [72]. The reason that these balls last twice as long is the inner core of the balls, coated with a nanocomposite coating that inhibits the permeation of air by 200% from the inside the ball. The coating was developed by InMat and utilizes aligned vermiculite platelets to impede the path of air from escaping. The platelets have an aspect ratio of 10000 to 1 and
significantly increase the path/travel distance for the air. The patented coating is known as Air D-fense® and is mixed at low viscosity and hence low shear stress. The technology is being extended to the tire industry, where these coatings would inhibit the migration of oxygen inside the tire, which affects the life of the steel tire cord. This further offers the opportunity to reduce weights, improve fuel efficiency and improve pressure retention and reduce recycling and incineration costs. InMat plans to continue the development of soccer balls, footballs, bicycle tires, automobile tires and truck tires based on this technology.

Nanocomposite coatings [68] based on UV curing of clay or colloidal silica filled acrylic or epoxy resins have also been studied. The coating had improved impact resistance, tensile strength, and hydrophobicity. Clay also imparts surface roughness and makes it an efficient matting agent. Nylon 11 coatings filled with nano-sized silica and carbon black have been reported to give an improvement of 35% in scratch, 67% in wear resistance and 50% decrease in water vapor transmission rate through nano reinforced coatings compared to pure polymer coatings [73].

However, it is important to ensure that nanoparticles are incorporated into coatings for textiles or fibers without losing their superb nanoscopic properties. It is well known that due to extremely high surface area, nanoparticles take any opportunity to agglomerate so as to minimize the surface energy. The dispersion of nanoparticles in the solvent media is influenced by stirring speeds, temperature profiles and the manner and order of addition of different components. Therefore, surface engineering of nanoparticles not only aids its true dispersion at the nanolevel in the coating solution but also ideally suits the needs of specific applications, further enhancing the properties of polymer nanocomposite based coatings. The chemistry of polymer nanocomposite formulations can be optimized to achieve coatings with best performance. The synthesis of these nanoparticles in larger quantities has been attempted [74] and M/s Degussa Inc (http://www.degussa.com) have already commercialized the process and launched a range of nanomaterials/nanoparticles for incorporating in the coatings.

13.8 Nanotechnology based fiber modifications

13.8.1 Nanocomposite fibers

Polymer nanocomposites are the advanced new class of materials with an ultrafine dispersion of nanofillers or nanoparticles in a polymeric matrix, where at least one dimension of nanofillers is smaller than about 10 nm. The nanofillers may have only one dimension in nano range – lamellar (e.g. layered silicate clays), two dimensions in nano range – fibrillar (e.g. carbon
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nanotubes) or all the three dimensions in nano range – spherical (e.g. metals/metal oxide nanoparticles and POSS (polyhedral oligomeric silsesquioxanes)) [75, 76].

Nano- vs microcomposite

Polymer nanocomposites have recently gained a great deal of attention because of superior properties when compared to neat polymers or conventional composites. The properties include high modulus, increased strength, improved heat resistance, decreased gas permeability and flame retardance at very low loadings of <5 wt% of nanofillers [77, 78]. In conventional composites, the mechanism of property enhancement is well understood and generally follows the rule of mixtures. On the other hand in nanocomposites, property enhancement exceeds the theoretically predicted values giving unique combination of properties synergistically derived from both the nanomaterials as well as the polymer matrix system. The volume and influence of the interfacial interactions increases exponentially with decreasing filler/reinforcement size and thus forms an additional separate phase known as interphase, which is distinct from the dispersed and continuous phases and hence influences the composite properties to a much greater extent even at low nanofiller loading (<5%). Therefore their properties are far superior to conventional composites. The interest in polymer nanocomposites further arises from the fact that they are light weight as compared to conventional composites because of the low filler loadings; they are usually transparent as scattering is minimized because of the nanoscale dimension involved and are still processable in many different ways even with nanoscale fillers embedded in the polymer matrix. The major challenges in nanocomposites are however ascertaining a high degree of dispersion of nanomaterials in the polymeric resin during nanocomposite synthesis and processing. Since the advent of nylon 6/montmorillonite (MMT) nanocomposites developed by Toyota Motor Co. Japan [79], relentless efforts are being made globally to successfully extend this concept to almost all types of polymer matrices with a range of nanomaterials used as reinforcing entities. Polymer nanocomposites thus exploit the fascinating and useful properties of nanomaterials for a variety of structural and non-structural applications such as automotives and packaging industries, building and construction, electrical and electronics, sports and medical devices [80–82].

Polymer nanocomposites also offer tremendous potential when produced in fiber form and offer properties that leapfrog those of currently known commodity synthetic fibers. Nanocomposite fibers that contain nanoscale embedded rigid particles as reinforcements show improved high temperature mechanical property, thermal stability, useful optical, electri-
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cal, barrier or other functionality such as improved dyeability, flame retardance, antimicrobial property, etc. These novel biphasic nanocomposite fibers in which the dispersed phase is of a nanoscale dimension, will make a major impact in tire reinforcement, electro-optical devices and other applications such as medical textiles, protective clothing, etc. [83]. The work on spinning of nanocomposites started about seven years ago and several research groups across the world are exploring the synthesis, fiber processing, structure-property characterization and correlation and molecular modeling of these unique new composite fibers.

13.8.2 Nanomaterials

All the three major types of nanofillers have been incorporated in polymeric nanocomposite fibers, viz. layered silicates (MMT), carbon nanotubes and nanofibers, metal oxide nanoparticles (TiO$_2$, ZnO, SiO$_2$, etc.) and hybrid nanostructured materials such as POSS and are briefly described below:

*Nanoclay (layered silicates)*

Layered silicates particularly Montmorillonite (MMT) clays, both natural as well as synthetic (alternatively referred to as 2:1 layered aluminosilicates, phyllosilicates, clay minerals and smectites), are the most commonly used inorganic nanoelements in polymer nanocomposite research to date [84, 85]. The chemical structure is:

\[
\text{Montmorillonite} = \text{M}_x(\text{Al}_4-x\text{Mg}_x)\text{Si}_8\text{O}_{20}(\text{OH})_4
\]

\[
\text{M} = \text{monovalent cation; } x = \text{degree of isomorphous substitution (between 0.5 and 1.3)}
\]

Their crystal lattice consists of two-dimensional layers where a central octahedral sheet of alumina or magnesia is fused to two external silicate layers. Depending on the precise chemical composition of the clay, the sheets bear a charge on the surface and edges, this charge being balanced by counter ions, which reside in part in the interlayer spacing of the clay. Stacking of the layers by weak dipolar or Van der Waals forces lead to interlayer or galleries between the layers. The galleries are normally occupied by inorganic cations (Na$^+$, Ca$^{++}$) balancing the charge of the oxide layers. These cations are readily ion-exchanged with a wide variety of positively charged species. The number of exchangeable interlayer cations are also referred to as the cation exchange capacity (CEC). This is generally expressed as milliequivalents/100 g and ranges between 60 and 120 for relevant smectites. In layered silicates, the Van der Waals interlayer or
gallery containing charge compensating cations (M$^+$) separates covalently bonded oxide layers, 0.96 nm thick, formed by fusing two silica tetrahedral sheets with an edge shared octahedral sheet of either alumina or magnesia (Figure 13.13).

Pristine layered silicates usually contain hydrated Na$^+$ or K$^+$ ions. One important consequence of the charged nature of the clays is that they are generally highly hydrophilic species and therefore naturally incompatible with a wide range of polymer types. Therefore clay often must be chemically treated in order to make it organophilic. When the inorganic cations are exchanged by the organic cations, these are called organically modified layered silicates (OMLS). Generally this can be done by ion exchange reactions with cationic surfactants including primary, secondary, tertiary and quaternary alkyl ammonium or alkyl phosphonium cations. These cations in the organosilicate lower the surface energy of the inorganic host and improve wetting characteristics of the polymer matrix or in some cases initiate the polymerization of monomers to improve the strength of the interface between the inorganic and polymer matrix [86].
Carbon nanotubes (CNTs) were first reported by Ijima in 1991 [87]. Since their discovery, CNTs have been the focus of considerable research because of the unique and unprecedented mechanical, electrical and thermal properties they exhibit (Table 13.3).

**Table 13.3 Physical properties of carbon fiber, carbon nanofibers and carbon nanotubes**

<table>
<thead>
<tr>
<th>Serial no.</th>
<th>Properties</th>
<th>Carbon fibre¹</th>
<th>Carbon nanofiber²</th>
<th>Carbon nanotube³</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Diameter (nm)</td>
<td>5000</td>
<td>3000–20000</td>
<td>SWCNT (rope) –20</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>SWCNT (single) –1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MWCNT –30–40</td>
</tr>
<tr>
<td>2</td>
<td>Length (nm)</td>
<td>–</td>
<td>30000–100000</td>
<td>1000–5000</td>
</tr>
<tr>
<td>3</td>
<td>Modulus (GPa)</td>
<td>465</td>
<td>400</td>
<td>1090–1260</td>
</tr>
<tr>
<td>4</td>
<td>Tensile strength (GPa)</td>
<td>3.92</td>
<td>2.7</td>
<td>13–52</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>SWCNT (rope form)</td>
</tr>
<tr>
<td>5</td>
<td>Strain to break (%)</td>
<td>0.8</td>
<td>1.5</td>
<td>SWCNT – 5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MWCNT – 12</td>
</tr>
<tr>
<td>6</td>
<td>Density (gm/cc)</td>
<td>1.88</td>
<td>1.80</td>
<td>1.3–1.5</td>
</tr>
<tr>
<td>7</td>
<td>Thermal conductivity (Wm⁻¹K⁻¹)</td>
<td>1400 (Max. achievable)</td>
<td>20</td>
<td>1800–3000</td>
</tr>
<tr>
<td>8</td>
<td>Electrical resistivity (μΩ cm)</td>
<td>3000 (vapor grown)⁴</td>
<td>1000</td>
<td>10⁻⁴ at 300 K (metallic SWCNT)</td>
</tr>
</tbody>
</table>

**Sources**

Carbon nanotubes are long, slender fullerenes where the walls of the tube are hexagonal carbon (graphite structure) and often capped at each end by hemi-fullerenes. Carbon nanotubes can be visualized as a sheet of the graphite that is rolled into a tube (Figure 13.14). Unlike diamond, where a 3-D diamond cubic crystal structure is formed with each carbon (sp\(^3\) hybridized) atom having four nearest neighbors arranged in a tetrahedron, graphite is formed as a 2-D sheet of carbon (sp\(^2\) hybridized) atom arranged in a hexagonal array. In this case each carbon atom has three nearest neighbors [88].

Carbon nanotubes are basically of two kinds – singled-walled (SWCNT) and multi-walled (MWCNT). SWCNTs can be considered as single graphene sheet rolled up into a seamless cylinder where graphene is a monolayer of sp\(^2\) bonded carbon atoms. MWCNTs are multilayered and simply composed of concentrically arranged single walled carbon nanotubes with a central hollow core with interlayer separation of almost 0.34 nm, an indication of the interplane spacing of graphite. A special case of MWCNTs is double walled carbon nanotubes (DWCNTs) that consist of two concentric graphite cylinders. DWCNTs are expected to exhibit higher flexural modulus than SWCNTs due to two walls and higher toughness than regular SWCNTs due to their smaller size.
The properties of nanotubes depend on atomic arrangement (how the sheets are rolled), the diameter and length of the tubes and the morphology of the nano-structure. The graphene sheets can be rolled up into tubes in various ways and are described by the tube chirality (or helical or wrapping), which is defined by the chiral vector \((C_n)\) and the chiral angle \((\theta)\):

\[
C_n = n \mathbf{a}_1 + m \mathbf{a}_2
\]

Where the integers \((n, m)\) are the numbers of steps along the unit vectors \((a_1\) and \(a_2\) of the hexagonal lattice. Using \(m\) and \(n\), the three different types of orientation of the carbon atom around the nanotube circumference is specified as armchair, \((n = m)\), zigzag \((n = 0, m = 0)\) or chiral \((\text{all others})\) (Figure 13.15). The chirality of the carbon nanotubes has significant implications on its properties, especially on the electronic properties. All armchair SWCNTs are metallic with a band gap of 0 eV. SWCNT with \(n - m = 3i\) (i being an integer \(\neq 0\)) are semi-metallic with a band gap of few MeV, while SWCNT with \(n - m \neq 3i\) are semiconductors with a band gap of 0.5–1 eV. MWCNTs contain a variety of tube chiralities, so that their properties are more complicated to predict.

MWCNTs and SWCNTs are mainly produced by three techniques: arc discharge, laser ablation and chemical vapor deposition (CVD). The various aspects of nanotube production, purification, suspension, filling, functionalization and application as well as the fabrication and characterization of polymer nanocomposites with various types of
nanotubes have been extensively reported in literature [89]. All known preparations of the carbon nanotubes give mixtures of nanotube chiralities, diameters and lengths along with different amounts and types of impurities. This CNT heterogeneity makes purifications of carbon nanotubes an essential requirement before they can be put into any application including synthesis of polymer/CNT nanocomposites.

CNTs exhibit exceptional material properties that are a consequence of their symmetric cage-like structure and exceed those of any previously existing materials. The high aspect ratio of CNTs coupled with a strong intrinsic Van der Waals force of attraction between nanotubes combine to produce ropes and bundles of CNT. SWCNTs where the attractive force is very high (0.5 eV per nm of CNT to CNT contact) are more prone to form ropes and bundles. Ropes refer to collections of SWNTs and are more uniform in diameter to form hexagonal lattices, while bundles are non-crystalline collections of SWCNTs or MWCNTs. It is for this reason that dispersion of CNTs in solvent to produce suspensions, need assistance of either sonication or surfactant addition or both combined besides mechanical stirring.

POSS

Polyhedral oligomeric silsesquioxanes (POSS) are an important class of nanostructured hybrid inorganic-organic materials and are widely regarded as one of the most promising and rapidly emerging nanocomposite materials. POSS molecules are of nanosized polyhedral or cage-type structure consisting of an inner silicon-oxygen (\(\text{SiO}_{1.5}\)) based framework with organic substituents (R) on outer corners and with a precisely defined silicon-silicon distance of 0.53 nm (Figure 13.16). The ratio of Si:O is 2:3.

The average diameter of a POSS molecule is 1.5 nm. The inorganic silicon-oxygen core is thermally and chemically robust. The organic substituents (R) are either totally hydrocarbon in nature or a range of polar structures and functional groups are also included. These organic substituents can be totally reactive or nonreactive or a combination of the two, depending on the synthetic design (Figure 13.16). According to the nature of the organic substituents, they undergo nucleophilic substitution or hydrosilylation. The density of POSS chemicals is typically in the range of 1.12 gm/ml, despite a high molar mass of about 1000 amu. The reason is most of the volume is taken up by the organic (R) groups, which lie on the outside of the cage whereas the core occupies only 5% of the overall cage volume. The physical state (solid, wax or liquid) ranges from low solubility...

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Various types of POSS structure [90].

system with $T_m \sim 400^\circ C$ to wax or grease or low temperature flowable oil depending on the nature of the organic (R) groups along with the topology of POSS cages [90].

POSS was first developed by the US Air Force for aerospace applications. POSS originally cost $5000$ per pound (454 g) and took up to three years to produce. Due to simplified and redesigned process chemistry, now
production cost has come down but still it is expensive. A variety of POSS monomers in form of solids or oils are now commercially available from Hybrid Plastics Company Co. (http://www.hybridplastics.com/), USA. A wide range of POSS compounds is now available that contain a combination of covalently bonded reactive functional groups that make them suitable for polymerization, surface bonding or other chemical transformations [91]. POSS moieties can thus be easily incorporated via copolymerization, grafting or blending into common plastics. POSS incorporated polymeric materials show significant improvements in use temperatures, oxidation resistance, surface hardening and enhanced mechanical properties. Reductions in flammability, heat evolution and viscosity during processing are also added advantages [92].

### 13.9 Polymer/clay nanocomposite fibers (PCNF)

Incorporation of organoclay into a polymer matrix has been used to modify the fiber and textile performance. The main effects are in physical properties, e.g. enhanced tensile modulus and strength, reduced thermal shrinkage, controlled electrostatic behavior, high storage and reduced loss modulus under dynamic mechanical testing conditions. In addition, presence of hydrophilic clay also enhances moisture absorption (at a slow rate), dyeability, biodegradability and chemical resistance. Since clay platelets block UV irradiation PNCFs exhibit improved weatherability. Reduced flammability of PNCFs arises due to inherent nature of clay to reduce the burning rate and form char. With these improved properties, these fibers have an edge over their neat counterparts in applications of both domestic and industrial textiles [83, 93–97].

PCNFs have been mostly spun through three basic methods of fiber spinning [83]:

- Melt spinning
- Solution spinning
- Electrospinning.

The major challenges here are achieving maximum extent of intercalation and exfoliation of clays in the polymer matrix (Figure 13.17). Owing to hydrophilic character of clay, the preparation of PCNF is easiest with water-soluble polymers. The increasing nonpolarity and hydrophobic nature of polymers, makes it more difficult to disperse the hydrophilic clay in an organic matrix. The strategy used to overcome this problem is two-fold:

- Modification of clay so as to make it organophilic
- Use of compatibilizer.
Organomodified clays have a quaternary ammonium ion with paraffin substituents, e.g. dimethyl di-hydrogenated tallow ammonium chloride (2M2HT) intercalated in interlayer galleries replacing the Na\(^+\), K\(^+\) cations making the clays organophilic and thus more dispersible in the polymer system. However, there is a tendency of quaternary ammonium ions to degrade at the higher polymer processing temperatures, i.e. 150–180°C. Attempts have been made to have groups such as 1,2-dimethyl-hexadecyl-imidazulam-MMT (IMD-MMT) and didecyl–triphenyl phosphonium-MMT (Cl2 PPH-MMT) which have higher thermal stability [98].
Nanoclay reinforced fibers have been produced from a number of polymers but only PNCFs based on polyester (PET) and polyamide (nylon 6 and nylon 66) are being reviewed in the section below.

13.9.1 Polyamide/clay nanocomposite fibers

Polyamide-6 (PA 6)/clay nanocomposites have been widely investigated due to their much superior tensile strength and modulus, improved heat resistance (heat distortion temperature increases from 65°C to 120°C) as well as excellent gas and water barrier properties [99–100] over their neat PA6 counterparts. However, specific studies on spinning and characterization of polyamide/clay nanocomposite fibers have been relatively few and are reviewed here [101–112].

Nylon 6 (PA 6)/clay nanocomposite fibers (PAMC) are generally produced using the melt-spinning route, where nylon 6 chips are compounded with MMT nanoclays during the melt extrusion process using mostly twin-screw configuration. Attempts have also been made to spin fibers of PA 6/clay by in-situ polymerization technique using double screw reactive extrusion [103]. The molecular weight and weight distribution measured for the polymerized PAMC fibers measured by GPC showed a tendency to first increase and then decrease with increasing screw speed. It succeeded in inserting a covalent linkage between OH-groups of organomodified silicate and the amino groups on PA 6.

The compounding of MMT clays into polymeric matrix is usually done on a single or double screw extruder, where the dispersive forces, i.e. shearing action inside the barrel, is responsible for breaking the minor components domain, i.e. clay particle to desired size and get homogeneously distributed in the matrix (Figure 13.18). It is reported that extensional flow mixer generates better dispersion and distribution mixing and is claimed to be orders of magnitude more efficient than shear mixing [83]. Hence the extensional flow mixer has been designed as an inexpensive mixer to be attached to either single or twin-screw extruder.

PA 6/clay nanocomposite fibers have also been produced using a high-speed melt spinning process at take-up velocities of 1–5 km/min. The clay used in all these researches is the most widely used organomodified MMT clay at the 2–5-wt% loading. Careful control of drawing conditions such as drawing temperature and ratio and subsequent annealing step are important to get the desired properties and has been investigated by several researchers [102, 104, 106, 110, 117]. The effect of clay on the fiber structure development and resultant properties has been reported by several researchers [101, 103–104] but more in detail by Giza et al. [110, 111] from Tokyo Institute of Technology, Japan.
PA 6 generally crystallizes in two crystalline forms, α form, which is monoclinic, and metastable γ form, which is pseudohexagonal. The α form generally grows by slow cooling from the melt or develops on annealing or drawing. The γ form is obtained by fast cooling from the melt notably in fibers spun at moderate spinning rates. The nanoclays in PA 6/clay nanocomposite fibres are reported to behave as γ-nucleating agents in both injection molded samples as well as oriented spun fibers [112]. However, these rapidly turn into alpha crystal form upon drawing. Matrix shearing in between the MMT platelets during drawing imparts better packing and cohesion to hydrogen bonded planes and thus improves their thermal stability as compared to those of neat PA 6 fibers [112]. PA 6/clay composite fibers also showed higher crystallinity in the case of whole take up range (1–5 km/min) as compared to neat PA 6 fibers (1–7 km/min) although the clay content does not affect the crystallinity values when produced using high speed spinning much. The PA 6/clay fibres when spun at such high

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speeds show better orientation of crystalline entities and also higher biefringence values, but only in the low take up region (up to 3 km/min). The orientation induced crystallization led to direct formation of \( \alpha \) form crystals in the spin line, which started at 2 km/min for PA 6/clay fiber and at 4 km/min for neat PA 6 fiber as revealed by WAXD studies. The PA 6/clay nanocomposite fibers showed superior Young’s modulus at all take up velocities but tenacity was higher only in low temperature velocity region where crystallization in the spin line of neat PA 6 did not occur yet [110, 111]. Thus nylon 6/clay nanocomposite fibers show reduced crystallinity and lower molecular orientation, which explains the observation of no significant improvements in tenacity in most of the studies. However, stiffness of intercrystalline regions and role of rigid clay particles in stress transmission behavior explains the superior modulus in PA 6/clay nanocomposite fibers.

Apart from improved mechanical properties the presence of clay in PA 6/clay fibers also affects its dyeing ability. The DSC studies show that presence of clay in PA 6 induces \( \gamma \) crystalline form, increases the crystallization temperature and decrease the melting point. A higher amorphous content observed improves the accessibility for disperse dyes. Thus PA 6/clay yarn dyes itself faster with disperse dyes than an unfilled PA 6 yarn while it gives opposite effect with acid and metal complex dyes. In both cases, nanoclay fixes on the amino sites and thus prevents fixation of acid or metal complex dyes [103]. The incorporation of clay in PA 6 fibers also leads to improved fire resistance and lowered flammability properties, thus offering a new promising route for flame retardant textiles with permanent effect and at low cost [108, 109].

13.9.2 Polyester/clay based nanocomposite fibers

There are significant opportunities for enhancing the properties of polyester fibers by incorporation of inorganic layered silicate nanoclays (MMT), as revealed by a number of research studies and patents reported in the literature [113–119]. Poly (ethylene terephthalate), PET/MMT clay nanocomposites are generally prepared by in situ interlayer polymerization [113–115] and then melt spun to produce monofilaments with varying draw ratios. The nanocomposite is prepared by mixing ethylene glycol, dimethyl terephthalate in the presence of organically modified silicate (0.5–15%), 0.001–1.0% additives and/or 0.001–1.0% catalyst (oxide acetate of Sb, Ge, Ti or Sn) and then polymerizing it [114]. The organic modifiers used for MMT clays are mostly phosphonium cationic salt [94, 113], which make the organomodified clays thermally stable and thus able to withstand the high temperatures involved during PET polymerization. The clay is generally well dispersed in the PET matrix as observed under scanning (SEM)
and transmission electron microscopes (TEM), but some clays were agglomerated at a size level in the range 10–20 nm [116]. The interlayer distance of MMT dispersed in the nanocomposite fiber was further enhanced because of the strong shear stresses during melt spinning [117].

The structure and properties of the PET/clay nanocomposite fibers have been studied as a function of organoclay content and the draw ratio (DR) given. The PET/clay hybrid nanocomposite fibers generally show improved thermal stability and mechanical properties as compared to their neat counterparts [116] even at low organoclay contents (<5 wt.%). The composite fibers also exhibit improved dimensional stability, higher modulus and low shrinkage. The tensile properties generally increase with increasing clay content up to DR = 1. However the values decrease with increasing draw ratio (DR = 1–6) and also higher clay content beyond a critical concentration [94, 118]. DSC and WAXD test results show that incorporation of clay generally accelerates the crystallization of PET but the crystallinity and orientation of drawn fibers is generally lower than that of drawn neat PET fibers [95]. The strong interaction between MMT and PET restricted the motion of PET chains, which developed a special continuous network structure and inhibits thermal shrinkage of PET and also improves other thermo-mechanical properties of PET fibers.

The PET/clay nanocomposite fibers with improved properties have the potential of finding application in tire cords to reinforce the rubber [96, 119]. High strength, heat resistance, heat resistant adhesion and low heat emission properties are required for tire and tire cords. Polyester tire cords are more attractive for this use but have some weaknesses. PET/clay nanocomposite fibers can overcome some of the weaknesses of the neat PET fibers and thus enhance the stability, uniformity and ride comfort. The incorporation of clay in polyester fiber also improves its dyeability with disperse dyes and making deep dyeable polyester a possibility [114].

13.10 Carbon nanotube (CNT) based nanocomposite fibers

Nanocomposites based on polymers dispersed with carbon nanotubes and nanofibers are beginning to receive substantial attention today because the carbon nanotubes offer immense potential to impart unique mechanical, electrical and thermal properties to polymeric fibers both polyester and polyamides [120]. Carbon nanotubes are characterized by high flexibility, low mass density and large aspect ratio, i.e. 300–1000. Some nanotubes are stronger than steel, lighter than aluminum and more conductive than copper. The theoretical and experimental results on single wall carbon nanotubes (SWNTs) show extremely high tensile modulus (640 GPa to
1 TPa), tensile strength (150–180 GPa) and failure strains too, of the order of 5 to 20%. SWNTs also possess very high electrical and thermal conductivity (theoretically >6000 W/m·K). Depending on their structural parameters, SWNTs can be metallic or semi-conducting [120]. The first nanocomposites based on CNTs were reported by Ajayan et al. in 1994 [121, 122]. The major challenges in polymer/CNT nanocomposites are as follows.

The ‘less than ideal’ and inconsistent nature of CNTs: all known methods of CNT production, give a mixture of CNTs with different chiralities, diameters and lengths, along with varying amount of impurities and structural defects within the sample or that from different batches. Hence, it is very difficult to obtain reproducible results [120]. Dispersion of CNTs in a polymer matrix is also very crucial to achieving the predicted properties. Due to Van der Waals interaction, SWNTs typically form bundles of 30 nm in diameter containing several hundred tubes within a bundle. These tubes are also highly entangled and the degree of entanglement depends on their length. Various physical and chemical approaches have been tried to disperse them, which include sonication, choosing the correct dispersion medium and use of surfactants to aid in dispersion [123]. Another great challenge is the efficient translation of CNT properties into the polymer matrix, which means ensuring a good interfacial interaction between the two; CNTs being inert, are not easily dispersed and will not interact with most of polymer matrix systems.

The approaches used are to functionalize the CNTs by oxidation, end cap functionalization with long aliphatic amines and sidewall functionalization using fluorination, alkylation, etc. [124]. Polymer grafting to create functional groups on CNTs has also been reported, which improves their dispersion as well as the interfacial interaction [125]. Non-covalent functionalization is achieved by adsorbing different polymers onto SWNT to improve dispersion. The SWNT solubilization by ‘polymer wrapping’ is also an alternative method for tuning the interfacial properties of nanotubes [126] (Figure 13.14).

Carbon nanotube orientation preferentially along the fiber axis plays a very critical role in imparting high tensile properties in CNT based polymeric nanocomposite fibers. SWNTs with a density of 1.3 g/cm³ have the potential of producing a high performance fiber with at least twice the specific tensile strength and specific tensile modulus of the presently known state-of-the-art PAN based carbon fiber [120, 123, 127] even at very low loading. However, large scale economic synthesis of CNTs and the ability to synthesize CNTs of uniform diameter and chirality will be critical to the commercial success of this new advanced class of materials for both molded parts, films as well as fiber forms. The CNT based polyester and polyamide nanocomposite fibers will be reviewed in this section.
13.10.1 Polyester/CNT based nanocomposite fibers

Oriented polymer systems such as fibers provide a unique opportunity to align the CNTs in one particular direction during the process of melt spinning and the subsequent drawing step. However one of the obstacles to widespread use of application of the nanotubes is the inability to orient them in one particular direction. Jose and coworkers at University of Alabama at Birmingham [128], focused on aligning carbon nanotubes in Vectra (a thermotropic liquid crystalline polymer) as well as polypropylene matrix during melt spinning into fibers. CNTs in two different weight percentages (0.5% and 1.0%) were used. Correlation of CNT alignment along the fiber axis and dispersion in the polymer matrix was studied using TEM (Transmission Electron Microscopy). Significant improvement in mechanical properties, i.e. tensile strength and modulus and a substantial increase in onset of decomposition temperature indicated both a good dispersion as well as a highly aligned CNT system.

Inherent hierarchical structure was observed in the form of fibrils that develop in vapor grown carbon nanofibers (VGCF) dispersed in a liquid crystal polymer matrix by Rohatgi et al. [129]. The tensile properties were sensitive to the diameter of extruded filament, the amount of VGCF added and other parameters related to the extrusion process. There is a noticeable increase in mechanical properties of composite filaments with decreasing filament diameter irrespective of VGCF content.

13.10.2 Polyamide/CNT nanocomposite fibers

There are several reports on nylon/CNT nanocomposites, where CNTs (single or multiwalled) or CNFs (carbon nanofibers) are infused into the polymer through liquid route using sonication or dry route followed by melt mixing in a single or twin screw extruder [130]. Alignment of CNTs and CNFs in the compositions was enforced during extrusion or subsequent stretching process. However, fullest potential of CNT reinforcement has not been harnessed in polyamide/CNT nanocomposite fibers, primarily because of lack of alignment or failure to develop strong interfacial bonding between CNTs and polymer.

Mahfug and co-workers [131] report a method to fabricate nylon 6/CNT reinforcement, where the MWNT reinforcement at low loading (<1 wt%) leads to 50% increase in tensile strength which is almost twice that of unreinforced nylon and never reported elsewhere in the literature. The spectacular enhancement of mechanical properties is attributed to the successful alignment of CNTs and the high interfacial shear strength developed during the fabrication process. The methodology described by them is different from that generally reported and involves special distributive
mixing of CNTs using a specially designed die and a prolonged thermal stabilization process prior to melt extrusion. These polyamide/CNT nanocomposite fibers have a great potential for use as multifunctional textile materials in several technical and advanced composite applications [132].

### 13.11 Nanoparticle based nanocomposite fibers

Preparation of PA 6/nano titanium dioxide (TiO$_2$) composites and their spinnability has been reported by Zhu et al. [133] of Dong Hua University, Shanghai, China. The nanoscaled TiO$_2$ particles were surface treated with coupling agents prior to mixing with molten PA6. The composite fibers showed improved mechanical properties as compared to pure PA 6 fibers and composite fibers with unmodified TiO$_2$.

PET/SiO$_2$ nanocomposites were synthesized using in situ polymerization and melt spun into fibers. The fibers PET/SiO$_2$ nanocomposite [134, 135] fibers showed a greater degree of weight loss as compared to pure PET fibers when they were hydrolyzed using alkali treatment. Superfine structures such as cracks, craters and cavities introduced due to this facilitated deep dyeing of PET/SiO$_2$ nanocomposite fibers. PET/silica as spun nanocomposite fibers show improved tenacity, modulus and lower heat shrinkage [136]. The silica nanoparticles get well dispersed in PET in the range 10 nm with narrow distribution. On the other hand, PET/TiO$_2$ nanocomposite fibers have been prepared by in situ polycondensation and melt spinning [137]. Nanotitania were first treated with a coupling agent to introduce functional groups on surface of titania particles and thus aid in dispersion of nanoparticles homogeneously. The UV protection property of these fibers was much enhanced (UPF > 50) because of high refractive index and absorption of UV light by the rutile form of nanotitania incorporated. However, these fibers had slightly reduced tensile strength and elongation at break. PET nanocomposite fibers have been reported with a range of other nanoparticles also, i.e. TiO$_2$, SiO$_2$, ZnO and CaCO$_3$. The nanoparticles are treated with low surface tension additive before mixing with PET [138].

POSS based nanofillers have been incorporated in PET and polyamide fibers/filaments and show improved thermomechanical properties and better retention of modulus at higher temperatures [139]. These fibers can find application in automotive tire reinforcement [140, 141]. However, the extent of reinforcement was found to be variable in the case of PET systems, which can be either due to low interaction between PET and POSS causing processing problems related to some water generation or chemical transformation of the POSS particles themselves [141].
13.12 Nanofibers

Recently, there has been an increased interest in producing nanofibers that are submicron size in diameter. Typically conventional melt blown ultrafine fiber diameter ranges from 2000 to 5000 nm, whereas polymeric nanofibres ranges from 50 to 500 nm. Nanofibers are characterized by an extraordinarily high surface area per unit mass (for instance nanofibers with 100 nm in diameter have a specific surface of 1000 m²/g) high porosity and lightweight. These unique properties of nanofibres make them potential candidates for a wide range of applications such as filtration, barrier fabrics, protective clothing, wipes and biomedical applications such as scaffolds for tissue engineering [142].

13.12.1 Electrospinning

The manufacturing technique most commonly associated with polymeric nanofibers is electrospinning, which is fundamentally different from conventional fiber production techniques, and is based on electrostatic forces. The idea dates back at least 60 years, but recent interest has been renewed by the pioneering work done by Dr Darryl Reneker at the University of Akron [143], who demonstrated electrospinning for a wide variety of polymer solutions including rigid rod polymers. The basic theory of the nano-fiber spinning process and the parameters affecting the process, thermal and mechanical properties of electrospun nanofibers has been much reviewed [144–147].

Electrospinning is a process that produces continuous polymeric nanofibres (diameter in the submicron range) through an action of an external electric field imposed on a polymer solution or melt. In this technique, a polymer is dissolved in a solvent (polymer melts can also be used) and placed in a glass capillary, which is sealed at one end and a small opening in a necked down portion at other end. A high voltage potential (730 kV) is then applied between the polymer solution and a collector near end of the capillary tube (Figure 13.19). When a voltage is applied to the polymer solution, the electric force results which cause a jet of polymer solution flowing from a droplet tip to be drawn toward a grounded collector. After the jet flows away from droplet in a nearly straight line, it bends and whips into a complex path and other changes in shapes occur, during which electrical forces stretch and thin it at very large ratios (about 100000) in a short distance and in less than one second. After the solvent evaporates, solidified nanofibres are left, mostly in nonwoven web form on the collector. This nanofiber deposition cannot be seen by the naked eye and is generally viewed under the scanning electron microscope (SEM) (Figure 13.20).
13.19 Schematic diagram of electrospinning set-up [153].

13.20 SEM images of (a) electrospun nanofibrous web, (b) cotton yarn coated with nanofibrous web.
The electrospinning process can be considered as a variation of the electrospraying process. In the electrospraying process, the surface of a hemispherical liquid drop suspended in equilibrium at the end of the capillary will be distorted into a conical shape in the presence of an electrostatic field. A balancing of the responsive forces resulting from the induced charge distribution on the surface of the drop with the surface tension of the liquid causes this distortion. Once a critical voltage ($V_c$) is exceeded, a stable jet of liquid is ejected from the cone tip. The jet breaks into droplets as a result of surface tension in the case of low viscosity liquids. For high viscosity liquids the jet does not break up, but travels to the grounded target. The first case is known as electrospraying and is used in industries to obtain aerosols and when applied to polymer solution and melts (second case) it generates polymer nanofibres through the electrospinning route (Figure 13.21) [147]. The morphological studies indicate that electrospinning process does partially orient the molecules in fibers, although to date all nanofibers produced are without any control over forces during orientation and crystallization.

Most of the literature on electrospinning has explored a variety of polymer/solvent systems from which fibers can be produced. Only a few studies have addressed the processing/property relationships in electrospun fibers. The structure and property of electrospun nanofibers are predominantly determined by the synergistic effect of solution parameters and electrostatic forces. Processing parameters considered are solution concentration and viscosity effects, spinning atmosphere effects, accelerating voltage effects and tip to target distance [146, 147]. Solution viscosity has been found to influence fiber diameter, initiating droplet size and jet trajec-
Increasing solution viscosity has been associated with the production of large fiber diameters. Spinning atmosphere has been associated with the jet splaying phenomenon also reported by Renekar et al. [148]. Other processing variables, such as acceleration voltage, electrospinning current and tip to target distance have also been investigated and linked to fiber morphology and defect structures [149].

13.12.2 Nanofiber morphology: effect of process parameters

**Voltage dependence [147]**

As the accelerating voltage is increased, electrospinning current generally increases step wise, which reflects in increase in mass flow rate from the capillary tip to the grounded target, when all other variables such as surface tension and charge density are kept constant. This is generally related to changes in originating droplet shape, which is related to bead defects in electrospun fibers. At low voltages, the electrospun fibers have a cylindrical geometry and distinct decrease in bead defects. As voltage increases, the density of bead defects significantly increases.

**Concentration dependence [147]**

Solution surface tension and viscosity also play an important role in determining the range of concentrations from which continuous fibers can be obtained in electrospinning. At low viscosities ($\eta < 1$ poise), surface tension has dominant influence on fiber morphology and below a certain concentration drops will form instead of fibers. At high concentration ($\eta > 20$ poise), processing is inhibited by the inability to control and maintain the flow of a polymer solution to the tip of the needle by the cohesive nature of high viscosity solutions. Although the range of concentrations that produce fibers will obviously vary depending on the polymer/solvent system used, the forces of viscosity and surface tension will determine the upper and lower boundaries of the processing window, other variables being constant.

Buer and co-workers [146] further investigated the various electrospinning equipment design parameters such as controlled feed rate of polymer solution using a syringe pump, choice of capillary material, extraneous currents and varying collector geometries (static, moving screen or plate) and construction material, etc. on the quality of nanofibers produced. Electrospinning is thus a fast and simple process of producing a wide range of nanofibers. The collected webs of nanofibers usually contain fiber with varying diameters from 50 nm to 2 $\mu$m. Sometimes the main fiber breaks up into many smaller filaments, a process called ‘splaying’, which happens because of electrostatic instabilities under certain conditions. As com-
pared to melt blown nanofibers, electrospun fibers have a much narrower diameter distribution.

The path of electrospinning jet being a combination of bending, looping and spiraling generally leads to nanofibers in mostly a nonwoven web form with no preferred orientation. There have been attempts to produce electrospun ultrafine fibers in an aligned fashion by the use of a cylinder surface moving linearly. The fibers are taken up tightly in a circumferential manner resulting in a fair alignment. The other route is to dampen the jet and control the deposition of electrospun fibers on a target. A series of charged rings acts as an electrostatic lens and changes the shape of the electric field. Modification in collector designs such as moving planar surface [150] or a cylinder with a rotating high speed [151] and a collector with a gap [152] are other approaches to collect nanofibers in an aligned form. An interesting and unique observation has been reported by Beck Tan and coworkers [147], where they observed that electrostatic effects also influence the macroscale morphology of electrospun textiles and may result in the formation of heterogeneous or 3D structures. They also observed that residual charge left on electrospun fibers affects the way they organize themselves on nonwoven textile substrate. 3D honeycomb structures were observed by them under a microscope when using dilute solutions onto a wire screen (Figure 13.22). Rabolt and coworkers [153]

![3D structure of a nonwoven fiber mat electrospun from a 7-wt% solution of PEO in water onto a cloth substrate [147].](image)

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have created a highly micro- and nanostructured ‘porous’ morphology in electrospun nanofibre (Figure 13.23), thus increasing their range of application significantly. The pores vary from densely packed, well-formed nanopores to large flat pores. The increased surface area of electrospun fibres was due to highly volatile solvents used.

**Other techniques to produce nanofibers**

Although electrospinning has been most widely reported in producing nanofibers, the low production rates of this process are a major limitation in upscaling this technique, and thus make it mostly a laboratory curiosity. However, attempts have been made to speed up the process using multiple syringes [154] as well as using needleless spinning. The production rate of nanofibers through electrospinning is generally measured in grams per hour.

Nanofibre Inc. Aberdeen, North Carolina, USA, have developed a unique process of producing nanofibers through melt blowing using a molecular die [155]. The fibers produced are a mixture of both micron and sub-micron sized fibers. But it is a relatively inexpensive technique and can
produce nanofibers in large quantities at a cost of $10 per kg. Although it is a relatively inexpensive technique and can produce nanofibers in large quantities at a cost of US$10 per kg, the broad range of fiber diameters produced is a concern. If perfected this technique can certainly take nanofiber production to a commercial future because of higher production rates as compared to electrospinning.

The other technique is that based on making bicomponent fibers that split or dissolve. The most researched is the production of islands in the sea fiber using a standard spin/draw process. The production rate is 5 kg/h at a take up speed of 2500 m per minute. Both PET and PA 6 nanofibers of diameters approx (300 nm) can be produced using EVOH as sea polymer, and the ratio being 50/50. Unlike electrospinning and melt blown technique, the nanofibers produced are in a narrow range and the projected cost is 1 to $5 per kg. Another possible approach is the use of bicomponent fiber spinning to manufacture nanofibres by the splitting process. The number of segments needs to be greater than 16 and use a water-soluble polymer in a small ratio along with PET and PA 6 [156].

Toray Industries Inc. [157] have developed a new technology for producing multifilament nanofibers comprising monofilaments whose diameter is in nanorange. The technology is versatile not only in its ability to be applied to commodity polymers such as nylon or polyester, but also in that nanofibers can be manufactured using existing production equipment. They have succeeded in producing for the first time nylon nanofibers having uniform diameter in nanorange through optimization of rheological properties of polymers themselves. Since nylon nanofibers are of multifilament continuous structure, they can be easily fabricated into a wide array of products while the orientation and shape of the nanofibers can be easily controlled, enabling their application to be extended into various fields. The nylon nanofibers produced have about 1000 times larger surface area than conventional fibers, thus adsorption and adhesion properties and novel functions attributable to fiber surface are also more pronounced and promise a variety of applications as advanced materials. The moisture absorption of these nylon nanofibers is about two to three times higher than conventional nylon, which is comparable to cotton.

In another ongoing NTC project at Georgia Tech, USA [158], a unique method of producing nylon and polyester nanofibers using ‘extrusion polymerization’ through mesoporous silica and aluminum channels as nanoreactors is being investigated (Figure 13.24). Unlike electrospun nanofibers, which are neither strong nor stiff, the polymer molecules in these fibers are not oriented. The nanofibers produced using this new technique will possesses high mechanical properties as they have a high degree of crystallinity from extended chain crystals formed inside the nanochannels.
The impact of nanotechnology produced mostly through electrospinning from a wide range of polymeric materials [159] but other techniques as described above [160] also show a from polymer nanocomposites, incorporating nanoclays, CNTs and other also been reported based on MMT clay [162, 163, 164] as well as CNT as through electrospinning at varying CNT content [166]. These composite 13.24 (a, b, c) Alumina nanochannels as prepared (pore size ∼40 nm), (d, e, f) pore formation and widening (pore size ∼100 nm), (e, f) longitudinal structure of nanochannels for alumina [158]. Thus, new researches in producing nanofibers in a continuous and aligned mono- or multi-filament form with more uniform diameter and narrow range have a potential to bring nanofibers in the market place, at a reasonable price and acceptable production rates. Nanofibers have been produced mostly through electrospinning from a wide range of polymeric materials [159] but other techniques as described above [160] also show a lot of promise in the future success of nanofibers.

**Polymer nanocomposite based nanofibers**

Recently, electrospinning has also been extended to making nanofibers from polymer nanocomposites, incorporating nanooclays, CNTs and other nanoparticles and adding a new dimension to nanofibers. Fong et al. [161] were the first to report electrospun nanofibers (∼100 nm) based on nylon 6/exfoliated MMT clay using hexafluoro-iso-propanol (HFIP) as the solvent. The resulting nanofibers had highly aligned MMT layers (normal to fiber axis) and nylon 6 crystallites (layer normal/parallel to fiber axis). Other polyamide 6,6 or polyamide nanocomposite electrospun fibers have also been reported based on MMT clay [162, 163, 164] as well as CNT as nanofiller [165]. Polyester/CNT composite nanofibers have been produced through electrospinning at varying CNT content [166]. These composite fibers can be further used to manufacture fabrics, antistatic materials,
electromagnetic shielding materials, high performance separation medium, reinforcing materials, electrical and thermal conductivity materials, wave absorbing materials, etc.

Fibers and nanofibers of PA 6/MMT clay nanocomposites (diameter between 100 to 500 nm) can be collected as nonwoven fabrics or as aligned yarns. The electrospinning process resulted in highly aligned MMT layers (layer normally perpendicular to the fiber axis) and nylon 6 crystallites (layer normal, parallel to fiber axis) [165, 167]. The PA 6/clay nanofiber web deposited on a nonwoven substrate as a continuous coating increased the contact angle and time required for water penetration relative to uncoated substrate [164].

Functionalized nanofibers for advanced applications

Functionalized nanofibers can be defined as nanofibers with specific additives for imparting special functionalities and capabilities to nanofibers thus widening the scope of their applications. The materials that have been added are metal/metal oxides at a nanorange, biological materials such as enzymes, drugs, CNTs (single as well as multiwalled) as well as nanoclays [168]. These value added nanofibers can be used effectively for several high-end applications in filtration, chemical protective clothing, biomaterials, drug delivery and tissue engineering. These polymeric nanofibers can also find applications in chemical and process industries such as catalysts, physical and chemical adsorption processes, etc. and are excellently reviewed by Ramkumar et al. [169].

Donaldson Company Inc. USA is one of the pioneers in this field and has commercialized the production of nanofiber webs from electrospinning for a broad range of filtration applications [170]. Nanofibers provide dramatic increases in filtration efficiency at relatively small decreases in permeability (Figure 13.25). Nanofiber filter media can contain airborne contamination in the personal cabin of mining workers. However, the production process and quality control of electrospinning nanofiber webs present special challenges. The parameters which have to be carefully monitored are polymer and solvent selection, electrical field control, solvent evaporation and management and nanofiber web formation and characterization. Quality control of nanofiber web also requires novel techniques based on real time measurement about web consistency.

13.13 Future trends

Nanotechnology has emerged as the ‘key’ technology, which has revitalized material science and led to the development and evolution of a new range of improved materials including polymers and textiles. This overview on
the impact of nanotechnology on textiles with a particular reference to polyester and polyamides indicates a clear shift to nanomaterials as a new tool to improve properties and gain multifunctionalities. Organized nanostructures as exhibited by either fibers, nanocoatings, nanofinishing, nanofibers and nanocomposites seem to have the potential to revolutionize the textile industry with new functionality such as self-cleaning surfaces, conducting textiles, antimicrobial properties, controlled hydrophilicity or hydrophobicity, protection against fire, UV radiation, etc. without affecting the bulk properties of fibers and fabrics.

However there are many challenges in the development of these products, which need to be intensively researched so that the wide range of
applications envisaged can become a commercial reality. An excellent
dispersion and stabilization of the nanoparticles in the polymer matrix is
crucial to achieving the desired nano effects. The tendency to agglomerate
due to an extremely high surface area is the major problem facing the
effective incorporation of nanoadditives in coatings/finishing as well as in
nanocomposite preparation. Surface engineering of nanoparticles and
combining them with functional surface-active polymers can bring the
nanoparticles onto fibers/textiles without losing their superb, nanoscopic
properties. More and more companies have specialized in making stable
dispersions of ZnO, Ag, SiO₂, etc. through the surface engineering of
nanoparticles.

Nanocomposite fibers based on clay show a lot of promise with their
improved properties such as stiffness, strength, heat deflection tempera-
ture, weatherability, moisture and dye absorption, reduced shrinkage,
flammability, etc, over their neat counterparts. These enhancements are
achieved with a small increase in density and cost. The most popularly used
nanofillers are the natural layered silicate clays, i.e. MMT although syn-
thetic clays with well-controlled and reproducible geometry and composi-
tion also need to be further explored. Incorporation of only a few percent
of carbon nanotubes (CNTs) can significantly alter not only the mechanical
properties but also enhance electrical and thermal conductivity of fibers.
However dispersion and exfoliation of carbon nanotubes continue to be a
challenge. Large-scale economic synthesis of CNTs particularly MWCNTs
and SWCNTs with controlled diameter and chirality will largely determine
the future commercial viability of these products. Nanofibers character-
ized by high surface area offer promising applications as efficient filters,
scaffolds for tissue engineering, vehicles for controlled delivery of medi-
cines and pesticides and more recently for biosensor applications. However,
faster production of nanofibers with controlled diameter, uniformity and
aligned morphology will play a crucial role in their future commercial
success.

To conclude: nanotechnology definitely has the potential to bring
revolution in the field of textiles. Estimated production of different
nanotechnology based technical textiles by the year 2010 is illustrated in
Figure 13.26. There is however a word of caution and the issues of concern
are:

(a) Large scale production of nanoparticles and their cost
(b) Impact of uncontrolled release of nanoparticles in the environment
and their effect on human health and ecology widely covered under
the domain ‘nanotoxicology’
(c) Practical philosophy and ethics on the widespread use of nanotech-
nology based products.
The impact of nanotechnology

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Part III

Applications of fibrous polyesters and polyamides
14
Polyester and polyamide fibres – apparel applications

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14.1 Introduction

Fibres are the building blocks of textile materials and there is a strong correlation between the fibre properties and properties of resultant textile products. The major end use application of textile fibres is its use in apparel fabrics. For thousands of years, cotton, wool and silk have been the most common and popular natural fibres for apparel. Invention of synthetic fibres especially polyamide and polyester and fibre engineering can produce, modify and tailor fibres in ways far beyond the performance limits of fibres drawn from silkworm cocoon, grown in the fields, or spun from the fleece of animals.

Polyamide refers to family of polymers called linear polyamides made from petroleum. The generic name polyamide fibre has the same meaning as nylon fibre, but nylon fibre is used principally in countries that derive their fibre technology directly or indirectly from the USA, and polyamide in countries that derive their technology from Germany. These two generic names continued to retain identical meanings when they were jointly redefined by ISO in 1977 as ‘having in the chain recurring amide groups at least 85% of which are attached to aliphatic or cyclo-aliphatic groups’. This change was made in order to exclude the new aromatic polyamide (aramid) fibres such as Nomex and Kevlar that had very different properties and end uses. There are two common methods of making polyamide for fibre applications. In 1931, American chemist Wallace Carothers was working on research carried out in the laboratories of DuPont Company where he focused on the miracle fibre ‘polyamide 6,6’. In this approach, molecules with an acid (COOH) group on each end are reacted with molecules containing amine (NH₂) groups on each end. The resulting polyamide is named on the basis of the number of carbon atoms separating the two acid groups and the two amines. Thus polyamide 6,6 which is widely used for fibres is made from adipic acid and hexamethylene diamine. Polyamide 6 and polyamide 6,6 have different performance characteristics and
Polyamide 6,6 has highly dense structure whereas polyamide 6 is an open molecular structure as shown in Fig. 14.1.

In another approach, a compound containing an amine at one end and an acid at the other is polymerized to form a chain with repeating units of \(-\text{NH}–[\text{CH}_2]_n–\text{CO}–\)\(_x\). If \(n = 5\), the polyamide is referred to as polyamide 6. In both cases the polyamide is melt spun and drawn after cooling to give the desired properties for each intended use. By 1938, Paul Schlack of I. G. Farben Company in Germany, polymerized caprolactam and created polyamide 6. It was the first synthetic fibre to go into full scale production and the only one to do so prior to World War II. Today, polyamide is used worldwide second only to cotton and polyester.

Polyester is a polymer, which is produced from coal, air, water and petroleum products. Among the earliest commercial polyesters were the alkyd resins or glyptals, based typically upon reaction of glycerol and phthalic anhydride to form a highly branched and ultimately cross-linked polymer and used in paints.\(^1\) When Dr Carothers was carrying out his fundamental research on methods for making long chain molecules such as those which constitute polymers and resins, he discovered not only the polyamides as used for manufacture of nylon fibres but also polymer known as polyester from diols and dicarboxylic acids that were not fibre forming. The discovery of terylene and its manufacturing was pioneered in Britain. Actually it was first made by J. R. Whinfield while he was working with J. J. Dickson in the research laboratories of The Calico Printers’ Association Ltd in Manchester in 1941. High melting fibre forming polyester was synthesized from ethylene glycol and terephthalic acid.\(^2,3\) It was first commercially produced in 1952 by the DuPont Company in the United States. DuPont initially named the fibre as fibre V but then coined it as Dacron in 1951. Polyester fibres have outstanding characteristics like wrinkle resistance and springing back into its crisp smooth shape, stretch and shrinkage resistance, strong and soft hand, excellent pleat retention and easy care properties. It is next to cotton in worldwide use.

Polyester fibre has good easy care properties and polyamide fibres have better elastic properties. These yarns are then either woven or knitted to develop fabrics. There is a strong relationship between textile materials and its quality. Major works in improving the properties, use of various forms and modification of fibres through texturing and spinning processes as well as blending have contributed to the wide expansion of these fibres.
Then quality of textile materials is manifested in different ways such as aesthetic appeal, feel or hand, etc. It is basically judged by how a textile material feels when touched or handled. Judging of fabrics is done both by subjective and objective methods.4

Thanks to the superior performance of fabrics made from polyester and polyamides, these fibres have now reached a stage that they can outperform ‘the natural fibres’ and can be engineered according to the market needs.

### 14.2 Properties of polyesters and polyamides and their suitability for apparel applications

Today’s apparel fabric maximizes enjoyment and comfort in indoors as well as outdoors and in any kind of weather. Waterproof, windproof and breathable through different finishing processes are just a few of many properties in demand. Some of the important characteristics of polyamide and polyester fabrics are given in Table 14.1.

The polyamide yarns of 3.3 tex are used for hosiery and 4–22 tex for apparel. The texturing process is used to stimulate the properties of natural staple yarns of increased bulk with benefits of thermal insulation, cover, softness and fullness and moisture transport.

Stretch properties can be imparted to polyamide fibres using textured nylon in combination with elastane yarns. Certain desired properties can be built into polyester and polyamide fibres during production. External form and constitution of fibres confer on the fibre certain properties which

<table>
<thead>
<tr>
<th>Polyamide</th>
<th>Polyester</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lightweight and very strong</td>
<td>Strong, stretchable</td>
</tr>
<tr>
<td>Stretchable and drapes well</td>
<td>Resistant to crease and shrink</td>
</tr>
<tr>
<td>Durable</td>
<td>Durable</td>
</tr>
<tr>
<td>Easy to clean, as dirt does not cling,</td>
<td>Easily washed</td>
</tr>
<tr>
<td>dries quickly</td>
<td></td>
</tr>
<tr>
<td>Resistant to abrasion and chemicals</td>
<td>Resistant to abrasion and chemicals</td>
</tr>
<tr>
<td>Low in moisture absorbency</td>
<td>Resistant to mildew</td>
</tr>
<tr>
<td></td>
<td>Does not absorb moisture making it hot and</td>
</tr>
<tr>
<td></td>
<td>clammy when worn in hot temperatures</td>
</tr>
<tr>
<td>Filament yarns provide smooth, soft,</td>
<td>Not damaged by sunlight or weather</td>
</tr>
<tr>
<td>long-lasting fabrics</td>
<td>Dries quickly. It is crisp and resilient</td>
</tr>
<tr>
<td>Spun yarns lend fabrics light weight and warmth</td>
<td>when wet or dry</td>
</tr>
</tbody>
</table>

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are advantageous for certain end products. The following modifications are also employed in order to achieve the various properties of the final product:

- fibre cross-section and surface filament size reduction (microfibre, superdrawing, alkalization)
- fibre blends
- bicomponent fibres
- bulk/texture (texturing and high bulk yarns)
- twisting, intermingling mixed yarns (hetero yarns, mixtures)

The fibre manufacturing process, in combination with further processing and finishing, results in a definite fibre structure regarding molecular structure and order, from which arise the fibre properties and finally the fibre capability in use. In this way, the final properties of the finished product are determined in terms of fibre properties.

Cross-section and filament surface form strongly influence the properties of the final product. In the case of melt-spun staple and yarn, the producer can determine the cross-section and surface form, as well as decide on whether to produce solid or hollow filaments, whereas these are partly determined by the strong influence of spinning conditions in the case of solution-spun yarns. Once chosen, the filament cross-sectional shape and form determine many yarn properties, which, in turn, determine the optical, tactile, physiological and technological properties of the final article. There are further opportunities to modify the cross-section and surface shape of fibre or fabric during further processing (e.g. texturizing, alkalization) in order to change the properties of the final product, but these are less effective than a modification introduced in the production stage.

Different polyester and polyamide fibre cross-sections and surface characteristics are shown in Figs. 14.2 and 14.3.⁷

![Cross-sectional form](chart)

Profiled encompasses the 2-lobed, 3-lobed (trilobal), rectangular (round, 4-lobed), star shaped (n-lobed).

### 14.3 Different fibre types for apparel purpose

Polyamides and polyesters have large number of types for different application but even then only some important types are widely used (Table 14.2).
The polyester and polyamide filaments and fibres of different aesthetic and functional properties are made by choosing polymer spin condition and draw conditions. In the case of polyamides mostly polyamide 6 and polyamide 6,6 are used and this comprises over 90% of commercial polyamide fibre production. The preferred polyesters are PET, PPT and PBT. Some important trade names and producers of polyamide 6, polyamide 6,6 and polyester are given in Table 14.3.

Three important characteristics of apparel fabrics for certain end uses are:

- Flame retardant – fabrics usually have a coating or film lamination against flame which provides a high level of protection and heat retardancy.
- Wet weather – water-repellent finished fabric provides normal comfort in the foulest weather due to their breathable and windproof characteristics. At the same time the fabric provides thermal insulation by
holding insulating air layers in the garment, maintaining warmth in the cold weather.

- Anti-static – this is to protect from static charge in the environments and anti-static fabric is durable along with being serviceable. It withstands regular cleaning abreast of retaining effective functional performance.
14.3.1 HM–HT cotton type polyester staple fibres

PET staple fibres can be blended with many natural and man-made fibres, and for this purpose have to match in linear densities and in the initial modulus. The HM–HT-cotton type (high modulus, high tenacity) shows similar properties as fine cotton. The HM cotton type is meant for blending with cotton. The viscose type is suited for blending with rayon and/or viscose staple. In spite of the different course of tenacity and elongation for more than 10%, the wool type and the anti-pilling wool type are particularly well suited for blending with wool. These PET staple fibres (high modulus high tenacity cotton type) with 1–2.5 dtex per fibre are gaining importance because of their ideal blending properties with cotton.

14.3.2 Polybutylene terephthalate (PBT)

Since its introduction around 1968, the production now increases annually about 10–15%, with increased 1,4 butandiol (butylene glycol). Butandiol (BD) is primarily obtained from acetylene and formaldehyde. New processes start with ethylene via promoethanol or with propylene via allylacetate and the oxo process. Polycondensation follows the scheme or analogue starting with TPA and with water as a byproduct. During the polycondensation butandiol is also regained.

\[
\begin{align*}
\text{CH}_3\text{O} &\quad \text{O} \quad \text{O} \\
\text{DMT} &\quad \text{C} \quad \text{C} \quad \text{O}-\text{CH}_3 + \text{HO}-(\text{CH}_2)_{4}-\text{OH} \rightarrow \\
+ &\quad 
\begin{cases}
\text{O} &\quad \text{O} \\
\text{PBT} &\quad \text{C} \quad \text{C} \quad \text{O}-(\text{CH}_2)_{4}-\text{O}- \\
\end{cases} + \text{CH}_2\text{OH} \\
&\quad n + \text{Methanol}
\end{align*}
\]

Yarns have a low moisture regain (about 0.4%) and very good crimp stability – important for BCF (bulked continuous filament) yarns. Considering the sales value of the byproducts, PBT yarn cost is already lower than PA-6 yarn cost. Another application is the combination with PET in bicomponent yarns to achieve crimp or dye effects.

Fibre forms also influence the textile process and the product scheme which finally segregate the apparel sections according to fibre form (Table 14.4).

There are various forms of polyamide and polyester fibre depending on the application areas:9
• Polyamide tow fibre

• Polyamide staple fibre – if staple fibre has to be processed into yarn on the cotton system then the tow is cut to fibres of 40 mm or less. Staple fibres for knitwear in which polyamide blended with wool are cut to staple length in the range 60–150 mm.

• Polyamide and polyester filament – polyamide flat filament yarn having 15–400 denier is applicable to clothing, industrial and knitted fabrics and POY is used for texturizing purpose. Different drawtextured polyamide and polyester filament yarns having high count micro-multifilament yarns of 50 to 600 denier are used for high grade clothing and decoration fabrics like ladies’ wear, pants and jacket fabrics, stockinng and casual fabrics. Semi-drawn and fully drawn polyester, polyamide and polyamide/polyester conjugate air textured filament yarns having 160–1000 denier are commercially used for sports wear, beach shorts and top coats as they provide a hairy feeling as natural cotton with characteristics of light weight, high wear resistance and easy to clean features. Some commercial blend proportions of polyester and cotton for wide variety of shirts, trousers and vests are described in Table 14.5. Various finishes to improve the appearance and function of shirts are used.

### 14.4 Blends of polyamide and polyester

Blending is one important process which provides the desirable attributes for apparel as single fibre cannot provide all desirable fabric properties.
Table 14.5 Polyester/cotton blend proportion for different types of apparel\(^{10}\)

<table>
<thead>
<tr>
<th>Type of apparel</th>
<th>Fabric</th>
<th>Blend</th>
<th>Finish &amp; care</th>
</tr>
</thead>
<tbody>
<tr>
<td>Casual shirt</td>
<td>106.3 g/m(^2), Dobby weave</td>
<td>60% cotton/40% polyester</td>
<td>Light soil wash</td>
</tr>
<tr>
<td>Performance polyester</td>
<td>134.6 g/m(^2), Poplin</td>
<td>100% spun polyester</td>
<td>Superior colour retention, Soft hand, Wickable finish, No-pill finish, Pre-cure durable press, soil release and wickable finish</td>
</tr>
<tr>
<td>Industrial work shirt</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Women’s work motion blouse</td>
<td>127.5 g/m(^2), Stretch poplin</td>
<td>75% polyester/25% cotton</td>
<td>Pre-cure durable press w/soil release and wickable finish, Industrial wash</td>
</tr>
<tr>
<td>Industrial work shirt</td>
<td>120.48 g/m(^2), Poplin</td>
<td>65% polyester/35% cotton</td>
<td>Pre-cure durable press w/soil release and wickable finish, Industrial wash</td>
</tr>
<tr>
<td>Geometric micro-check work shirt</td>
<td>110.5 g/m(^2), Poplin</td>
<td>65% polyester/35% cotton</td>
<td>Pre-cure durable press w/soil release and wickable finish, Industrial wash</td>
</tr>
<tr>
<td>Western style uniform shirt</td>
<td>113.39 g/m(^2), Chambray</td>
<td>65% polyester/35% cotton</td>
<td>Pre-cure durable press, Industrial wash</td>
</tr>
<tr>
<td>Performance knitted shirt</td>
<td>155.9 g/m(^2), Pique knit</td>
<td>50% polyester/50% cotton</td>
<td>Soil release, wickable finish, Industrial wash</td>
</tr>
<tr>
<td>Formerly work pants</td>
<td>240.97 g/m(^2), No-iron twill</td>
<td>65% polyester/35% cotton</td>
<td>Soil release, Home wash</td>
</tr>
<tr>
<td>Insulated vest</td>
<td>255.14 g/m(^2), Blended duck</td>
<td>65% polyester/35% cotton</td>
<td>Industrial wash</td>
</tr>
</tbody>
</table>

\(^{10}\) Polyester and polyamide fibres – apparel applications

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Blending helps use the positive attributes of both synthetic fibre like wear and easy care properties with better feel and high moisture absorbency property of natural fibres. Different natural and manmade fibres are blended in different combinations to produce a wide variety of yarns.

Polyester fibre can be suitably blended with cotton and viscose because of the complementary nature of the properties. The blend of polyester with viscose/cotton results in reduction of most of the negative features of polyester like poor moisture absorption, poor static dissipation, poor moisture vapour transmission, poor feel, warm, crisp hand, lower comfort, non-biodegradable, while negative features of viscose/cotton such as poor crease retention, poor wrinkle recovery, poor tenacity, lower abrasion resistance, higher staining tendency, difficulty of washing, lower colour fastness are overcome by the presence of polyester in the blend.

In general, the main motives behind blending are:

- Combination of merits of different fibres
- Reduction of cost by use of cheaper fibre with higher cost fibre in a blend
- Opportunity to produce colour effects.

Polyester and polyamide fibres which have a number of positive attributes when strength, elasticity and durability are considered, have a number of negative attributes as far as handle and comfort related properties are concerned. Comfort attributes of fibres like cotton and viscose can be combined with hard wearing and heat settable attributes of synthetic fibres to produce fabrics with the right balance of properties. Desirable or undesirable attributes of fibres can be affected by the actual textile construction. Yarn and fabric parameters, together with finishing treatments and garment design can considerably change the comfort level produced by a garment.

Synthetic-, chemical- and natural fibres each have their own specific properties, none of which alone may satisfy the requirement of a specific end use. In such cases, it can be advantageous to combine the properties of two or three fibre types in order to obtain an optimum for the end use in question or to enhance a particular product characteristic.

In principle, the following methods can be used to mix or blend fibres or yarns:

- Combining different staple fibres or staple fibre with natural fibre in flock or at carding
- Combining filament yarns and/or staple yarns by means of plying, twisting or entangling
- Combining yarns during fabric manufacture.

The following fibre combinations have proved themselves successful in terms of processability and optimization of product properties:
Polyester blends well with other fibres. It blends with wool, acetate, or rayon to improve the durability of the fabric and even to make it easy washable, if the percentage of polyester is high. It adds the quality of wrinkle resistance to the fabric and eliminates crushing of napped fabrics, and reduces fading. The fabric aggravates pilling problems after being blended with wool.

Polyamide blends well and blended fabrics have good dimensional stability, elastic recovery, shape retention, and abrasion resistance properties. Polyamide is very resilient, hence after blending it helps in eliminating the crushing of napped fabrics such as velvet.
14.5 Apparel applications of polyamide and polyester fabric

The application of polyamide expanded from the initial hosiery market to other apparel by exploiting easy care properties. In many textile applications, flat yarns with relatively coarse filaments were used in tightly woven or knitted fabrics. These fabrics had a limp and rather plastic handle, poor ability to wick moisture away from the body, and a strong tendency to build up static charges, leading to clinging and sparking. The fabrics were frequently shiny. In the 1970s the newly developed polyester/cotton blends offered superior comfort in many apparel applications and polyamide was displaced from these end-uses.

For apparel, producers engineer their products to meet the needs of particular uses – filaments have become finer and non-circular in cross-section, also textured yarns are used. These products, together with changes in fashion, led in the late 1990s to a revival of interest in polyamide for outerwear. The brand names like Tactel, Antron and Tactesse for particular applications are used.

14.5.1 Polyamide fabric

1. Polyamide is widely used in women’s hosiery and lingerie due to its high elongation and excellent elastic recovery quality. It is also used as sportswear, jackets, pants, skirts, raincoats, ski and snow apparel, windbreakers and children’s-wear.

2. Apparel: Blouses, dresses, foundation garments, hosiery, lingerie, underwear, raincoats, ski apparel, windbreakers, swimwear, and cycle wear.

14.5.2 Polyester fabric

1. Polyester clothing has a good stability and strength and is resistant to stretching and shrinkage. It is not damaged by sunlight or weather. It is widely used as dresses, blouses, jackets, separates, sportswear, suits, shirts, pants, rainwear, lingerie and children’s-wear.

2. Polyester fabric is manufactured in many weights and it is used as fibrefill in pillows and upholstery. In upholstery, polyester is generally blended with wool to eliminate crushing and reduce fading.

14.6 Comparison of polyesters and polyamides

Polyester and polyamide fibres both have attained their peak position in textile world and polyester is predicted to be dominating in the fibre race...
in coming years also. There are certain properties which separate the two fibres from most of the others and help to distinguish the area of application of polyester and polyamide fibres (Table 14.7).

**14.7 Pivotal fibre modification**

In recent years, polyester and polyamide variants such as stretch, microfibre, recycled, antimicrobial are increasingly used in addition to these fibres with traditional attributes.

Various innovative achievements through modification of technique or product have made it possible to confer the comfort and coolness of cotton, the soft warmth of wool and glamour and rustle of silk to polyester fibre. Cotton-like effects can be given to fine polyester staple of 1 denier or less crimped polyester filaments with irregular cross-section have improved moisture retention. The hollow polyester fibres have good thermal insulation. Thus there are multiple effects that are given to nylon and polyester. Some of the important modifications of polyester and polyamide fibres are discussed below.

### 14.7.1 Modification at polymer stage

ICI claims the preparation of improved linear polyester having better affinity for basic dyes by carrying out the polycondensation process in the presence of diphenyl sulphonie acids or their salts. ICI have also suggested a method of preparation of polyester of improved moisture regain by the addition of 5–10% by weight of sodium sulphate, of particle size less than 3 microns, to a slurry in ethylene glycol during polymerization.

<table>
<thead>
<tr>
<th>Properties</th>
<th>Polyester</th>
<th>Polyamide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial modulus</td>
<td>600–1200 gftex⁻¹</td>
<td>200–300 gftex⁻¹</td>
</tr>
<tr>
<td>Degree of elasticity</td>
<td>90–98 at 2% elongation</td>
<td>95–100 at 2% and 10% elongation</td>
</tr>
<tr>
<td>Moisture regain</td>
<td>0.4% at 65% RH</td>
<td>2.8–5.0%</td>
</tr>
<tr>
<td>Breaking elongation</td>
<td>12–55%</td>
<td>16–65%</td>
</tr>
<tr>
<td>Tenacity</td>
<td>25–54 gftex⁻¹</td>
<td>32–65 gftex⁻¹</td>
</tr>
<tr>
<td>Thermal conductivity</td>
<td>0.141 W/m K</td>
<td>0.243 W/m K</td>
</tr>
</tbody>
</table>
Lee *et al.*\textsuperscript{18} grafted acrylic acid by exposing impregnated drawn crystal-line polyester fibres to gamma radiation in the nitrogen atmosphere and found that the resulting fibre has good absorption and tensile properties and affinity for basic dyes. Hebeiish *et al.*\textsuperscript{17} have reported the properties of polyester/cotton blended fabric grafting with 2-methyl 5-vinyl pyridine using benzoyl peroxide as initiator. Fabric properties such as moisture regain, tensile strength and elongation at break have been found to increase with an increasing degree of drafting whereas crease recovery angle and electrical resistivity show a decreasing trend. It shows that certain desirable properties are adversely affected by attempts for making polyester hydrophilic. Blending turns out to be one of the important ways for imparting hydrophilic properties to the polyester apparels. And cotton/PET blends have been widely used in apparel applications because of improved comfort properties.

DuPont offers several stretch polyester derivatives based on its 3GT technology. The first 3GT product is named Sorona. Three carbon glycol is used to make these new fibres, unlike regular polyester which is made with two carbon glycol, hence the name 3GT. DuPont has developed a bio-source process that uses a fermentable sugar as a feed stock. It ferments sugar using a biocatalyst that converts the sugar into 1,3 propanediol (3G) which is purified to fibre grade. Properties of Sorona are:  \textsuperscript{18}

- Stretch and recovery are better than polyamide and regular polyester. Thus fabrics retain their shape and have enough stretch with recovery for comfortable movement
- Sorona, like polyamide, has lower modulus and is much softer
- It is easy to dye with same dyestuff as polyester without carrier.

Toray Industries, Inc. has developed a unique polyamide fabric Miramatte that realizes the softness and feel of polyamide and also features a high-quality plumage-like touch (‘Fine Feather Feeling’), with a refreshingly comfortable dry feel, and optimum drape. At the same time, Miramatte has excellent anti-ultraviolet and anti-transparent properties, and gives full play to its characteristics even if thin textile is used. The product is thus ideally suited to production of lightweight clothing.\textsuperscript{19}

Regarding the polyamide fibre to be used in Miramatte, Toray has increased the inorganic particles, which are effective for anti-ultraviolet and anti-transparent properties in the polyamide polymer, by approximately three times which is as much as the maximum density achieved with existing mass production. In addition, the particles are distributed uniformly not only in the fibre but also on its surface. It can be perfectly designed to develop cross-section variations and fine denier because of the uniform distribution of the inorganic particles at the stage of polymerization. In view of the consistent popularity of polyamide materials for sports
and casual wear in the fashion industry, Toray as the top producer of polyamide fibre in Japan has been carrying out research and development on new polyamide materials. The company plans to continue developing new polyamide products with features like:

- High-quality plumage-like touch
- Comfortable dry feel
- Optimum drape
- Refreshing feel
- Soft and matt colour tones with luxury feel
- Anti-ultraviolet property (UV covering ratio: more than 90%)
- Remarkable anti-transparent property
- Thin and lightweight clothing.

T 400 is polyester-based DuPont fibre developed with 3GT technology but it contains other components that produce low-to-moderate wovens, or true elastic stretch and recovery. It has stretch recovery, resistance to chemicals and abrasions are better than textured polyester and polyamide. It targets more 100% man made and blended fabric for ready-to-wear, typically with 20–30% Type 400 content.20

Nylstar inherited Elite stretch polyester from Schneer. Elite is modified polyester with permanent stretch properties and 98% recovery. The beauty of Elite is the ease with which it can bring staple spun yarns. The newest version of elite is a microfibre, which can be used for activewear.

14.7.2 Modification at spinning stage

Modification of polyester fibre – Ultra fine fibres

An ultra fine fibre is conventionally defined as a fibre of less than 0.7 denier. These fibres are often known as microfibres. An ultra fine fibre could be extruded by reducing the polymer output at the spinneret and drawing by a large drawing ratio.

Toray put the suede like new material Ecsaine on the market. Only ultra-fine polyester fibre can reproduce deer suede from the microscopic as well as macroscopic structural point of view. Ecsaine is made of homogeneous ultra-fine polyester fibres of less than 0.1 den and provides an epoch making appearance and hand as a clothing material. The pedal shaped conjugate fibre by DuPont was probably the first example of a potential ultra-fine filament.

As bending and torsional stiffness is inversely proportional to diameter, ultra-fine fibres are extremely flexible. The following characteristics can be shown:
• A softness, flexibility and smoothness
• A fine textile structure
• Formation of micro-pockets in fabrics
• A high filament density per textile surface
• Characteristic interfacial properties
• A small radius of curvature (resulting lustre and characteristics colour)
• A large aspect ratio (the ratio of length to diameter)
• A quick stress relief
• Low resistance against bending
• Fine edge.

The island in sea technology provides the industrial way of suede type artificial leather, silk like fabrics. The technology is further extended to spin the multi-component conjugate fibre, and a suede type artificial leather of high dyeability was developed with three component conjugate extrusion where the component (polyester and polyamide) ultra-fine fibres have sheath and core structure.\(^\text{21}\)

Water absorbent polyester fibres have recently been commercialized in Japan. Kilatt by Kanebo is a hollow fibre having a narrow slit through which water enters into the inner space and exits. Another water absorbent hollow fibre, Welkey, is made by Teijin.

14.7.3 Developments in blends

**Polyester and polyamide conductive fibre**

Apparel applications of conducting fibre are mainly in electrostatic discharge (ESD \(10^{-9}\) S/cm) applications in work-wear, dust-free garments. Typical bi-component fibres containing carbon black or white metal reach the requirement also in mixed fibre materials. However the conductive polymer materials reach 10–100 S/cm conductivity, and can be used in electromagnetic impulse (EMI \(10^{-3}\) S/cm) protection. Belltron commercial products have introduced carbon-black based sandwich structures having electric resistance \(10^8\) to \(10^{10}\) \(\Omega\)/cm and \(10^6–10^8\) \(\Omega\)/cm in 2.5 g/d tenacity polyester and polyamide fibre respectively.\(^\text{15}\) The comparable values for white metal powder construction were \(10^8–10^{10}\) \(\Omega\)/cm for both co-centric and side-by-side polyamide based bi-components 3 g/d tenacity fibre. The spinnability of the polymer melt has been shown to be a very complex challenge even for the homogeneous thermoplastic.

Dacron Plus is softer polyester with better drape. Dacron Duracotton, primarily for apparel, offers improved compatibility with cotton to create more durable fabrics that still have the look and hand of cotton but is enhanced by Dacron. DAK America’s Delcron brand, carried over from
the Akra side of the business, is offering a new moisture management product called Delcron Hydrotec with superior wicking properties and improved finished fabric costs. Delcron Colorbrite is a cationic dyeable fibre that achieves more brilliant colorations than typical polyester. Delcron high trek is a blend of almost equal parts of wool, acrylic and hollow polyester, created to be washable for garments as school uniforms and outdoor jackets.\(^2^2\)

KoSa recently introduced polyester yarns that are dyeable at lower temperatures than regular polyester, offering excellent printing and dyeing effects when blended with other heat-sensitive fibres such as spandex, wool and acetate. It is a single polyester yarn engineered with cotton like hand and appearance that also stretches. It is atmospherically dyeable and can be combined with other yarns for circular and seamless knitting, making it suitable for use in sports and thermal wear, intimate apparel, T-shirts and loungewear.

**Far infrared filament yarn**

This is a healthy care fibre with the property of heat and temperature maintenance and health protection. The yarn will be made from the polymer which is mixed with far infrared elements, so the far infrared filament yarn can emit electromagnetic wave of 4–14 μm. The wave can permeate into the deepest skin and hypodermic tissue and enable the water molecules inside the body to resonate. The water molecules violently vibrate and generate heat efficacy and then make the hypodermic deep temperature rise and capillaries expand. And it enables the water molecules of the cellular surface to be activated and speed the move of the calcium ion and therefore improve blood circulation and strengthen metabolism so that it vitalizes the tissue, quickens nutrition absorption and expels harmful substances in order for the body to acquire health and vitality. Its features:

- Vitalizing the skin with warmth-maintaining effects
- Prevention of chronic disease
- Increasing the immune system and health protection function
- Improving blood circulation and strengthening metabolism.

**Related end uses**

1. Suitable for applying to the fabric for metabolism improvement and warmth keeping
2. Functional underwear
3. Anti-varicosity hosiery
4. Exercise protector
5. Light and cold protective fabric and lining.
Anti-microbial filament yarn

Anti-microbial fibre is a fibre containing the inorganic anti-bacterial zeolite. The antibacterial zeolite is a multi-hole ceramic with silver. When the bacteria touch the ceramic, the silver ion and the enzyme in the bacteria cells will have a chemical action. It will restrain the activity of the bacteria and prevent the bacteria from procreating. Then it destroys the bacteria and achieves the effect of anti-bacteria and odour prevention. Related end uses are underwear, sweat shirts, hosiery, lining, cold protecting coats and gloves.

Polyamide filament trilobal bright

A trilobal cross-section makes good effect of light reflection and is used to produce clothing fabrics with high brightness, lustre and softness.

Polyamide/polyester conjugate micro-filament

Polyamide and polyester conjugates are usually made of 25% polyamide and 75% polyester. The yarn self splits in sanding when dyeing and finishing. So there is no need of caustic reduction, or only need of light weight reduction. This saves process cost, avoids waste water pollution and dyeing failure. The conjugate filaments have soft touch, lightness, graceful shine, and adequate bulkiness and are used to make high density fabrics with excellent wet penetration and ventilation properties, silk-like fabrics and artificial leathers. Excellent water absorption bath towel robe can be made from polyamide/polyester conjugate filaments.

14.8 Current market potentials

The development of man-made fibres in the last 40 years has been very great compared to natural fibres. Figure 14.4 illustrates the trend of man-made fibre growth in the past 10 years. The synthetic fibre segment produced a 3.2% higher output, mainly stimulated by 6.5% surging output of polyester filament yarns. The global polyester industry achieved a growth rate of 4.4% at 27.4 million tonnes. Much of the growth was driven by strong expansion of filament yarn that rose 6.5% to 16.3 million tonnes whereas the staple fibres showed slow growth of 1.6% to 11.1 million tonnes.

Asia accounts for 88% of the volume produced. Although several Asian countries suffered from reduced output volumes, this region has the ability to lift production by 6.4% to 24 million tonnes. Greater Europe and America ended the year with reduced production by 5.2% and 10.2% respectively.
China and India which produced an additional supply of 450 thousand tonnes and lower production rate in countries like Greater Europe, US, Taiwan and Brazil.

The steadily increased caprolactam and

14.4 Comparison of polyester and polyamide production with other fibres.

14.5 Production of synthetic fibres.

Filament fibre production has overtaken the staple fibre production in last 10 years (Fig. 14.5). Staple fibre business involves big players like PR China and India which produced an additional supply of 450 thousand tonnes and lower production rate in countries like Greater Europe, US, Taiwan and Brazil.

Polyamide fibres showed 1.7% increased output in 2006 by achieving 4.05 million tonnes (Fig. 14.6). The steadily increased caprolactam and
polymer prices, depressed housing market in the US and substitution to
other lower priced fibres slumped the staple fibre production by 8.0% to
376 thousand tonnes.\(^{22}\) The total filament market recovered by 2.8% to 3.7
million tonnes. Stagnant market conditions in Greater Europe and 4%
decline in America was facing a 8.6% increase in Asian production. PR
China expanded production by 21.5% to 825 thousand tonnes and Taiwan
witnessed a 3% increase in filament output. Meanwhile, the contribution
from Korean polyamide industry declined 6.5%.

### 14.9 Future prospects

According to the United Nations, world population will reach 6.82 billion
people by 2010. A world population of 6.57 billion corresponds to an
average per capita consumption of 10.5 kg and the increased response of
South East Asia in exports. The total man-made fibre production is esti-
mated to be 48.4 million tonnes in 2010.

In 1990 world polyester production was 8.58 million tonnes and polyam-
ide production was 3.88 million tonnes. By 2006, polyester production had
tripled to 27.307 million tonnes and polyamide reached 4.115 million
tonnes. And this increase forever changed the world map of fibre produc-
tion. Long the preserve of Europe and the United States, fibres now are
produced worldwide, with recent emphasis, particularly China, Taiwan,
South Korea and most recently, India. China’s share of the polyester fila-
ment production grew from 15.7% in 1990 to 62.8% in 2006.
It appears there will be a competition between India and China in polyester fibre manufacturing. Some big Indian textile and apparel companies have acquired world recognized brands for their polyester products. Figure 14.7 details the recent history of the major participants of polyester and polyamide and projects the volume by 2010.

Polyamide, the granddaddy of man made fibres, seems to be losing its share to polyester, overwhelmed by sheer volume if not performance. Many woven industrial and apparel fabrics seem to favour polyester. Polyamide’s dyeability is an advantage, but not sufficiently so to overcome the supply and variants available in polyester.

### 14.10 Acknowledgement

The help of Sangeeta Yadav in preparing the chapter is gratefully acknowledged.

### 14.11 References


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Polyester and nylon based textiles in biomedical engineering

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15.1 Introduction

One of the most innovative and growing aspects of the textile industry is the field of medicine, healthcare and hygiene. The extent of the growth of this domain is due to constant improvements and recent innovations in both textile technology and medical procedures. The first recorded use of fibres in medicine was mentioned in ‘Surgical Papyrus’ 4000 years ago. In the ‘Susanta Sambita’ of Indian Literature, written approximately 2500 years ago, a variety of materials including horse hair, leather strips and cotton are mentioned. With the passage of time, textiles have found their way into a variety of medical applications. In addition to protective medical apparel, textiles in fibre and fabric form are used as implants, filters and surgical dressings. Recent decades have witnessed major developments in medical textile production, the materials they are made of, and the technology used to produce them. Textile materials and products that have been engineered to meet particular needs are suitable for any medical and surgical application where a combination of strength, flexibility, and sometimes moisture and air permeability are required. Materials used include monofilament and multifilament yarns, woven, knitted, nonwoven fabrics, and composite structures. The numbers of applications are huge and diverse, ranging from a single thread suture to the complex composite structures for bone replacement, and from the simple cleaning wipe to the advanced barrier fabrics used in operating theaters [1]. Table 15.1 summarizes the main type of textile structures used for various healthcare and medical devices, including implants.

Medical textiles account for a huge market owing to the widespread need for them, not only in hospital, hygiene and healthcare sectors but also in hotels and other environments where hygiene is required. There has been a sharp increase in the use of natural as well as synthetic fibres in producing various medical products. The world medical devices market is represented by the Global Harmonization Task Force (GHTF). The GHTF...
Table 15.1 Fabric structure used in various healthcare and medical devices [2]

<table>
<thead>
<tr>
<th>Fabric structure</th>
<th>Uses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Woven</td>
<td>Gauze dressings, compression bandages, plasters, scaffolds, vascular prostheses, surgical gowns, drapes and hospital textiles such as sheets, blankets, pillowcases, uniform and operating room textiles, implants, knee supports and braces</td>
</tr>
<tr>
<td>Nonwoven</td>
<td>Surgical gowns, caps and masks, absorbent layers, fleeces, wipes, protective clothing, diapers, feminine hygiene products, incontinence products, wound dressings, scaffolds, implants, and antidecubitus fleece</td>
</tr>
<tr>
<td>Knitted</td>
<td>Compression bandages, vascular prostheses, stents, heart, valves, ligaments and tendons, surgical hosiery, blankets, wound dressings, stockings, elasticated net garments, pressure garments, finger bandages, flat bandages and spacer materials for knee braces, implants, nets and hammocks</td>
</tr>
<tr>
<td>Crochet</td>
<td>Compression bandages for compression therapy, cast cloth for orthopaedic casting bandages, wound dressings, bandages and implants</td>
</tr>
<tr>
<td>Embroidery</td>
<td>Implants</td>
</tr>
<tr>
<td>Braided</td>
<td>Sutures, soft tissue ligaments and implants</td>
</tr>
<tr>
<td>Composite materials</td>
<td>Diapers, feminine hygiene, incontinence products, wound dressings, scaffolds, implants and support systems for treatment of pressure ulcers</td>
</tr>
</tbody>
</table>

comprises representatives from five founding members (European Union, United States, Canada, Australia, and Japan). On the basis of DRA’s (David Rigby Associates) research, over 1.5 million tonnes of textile materials, with a value of US$ 5.4 bn, were consumed worldwide in the manufacture of medical and hygiene products in 2000. This is predicted to increase in volume terms by 4.5% per annum to 2010 to reach 2.4 million tonnes with a value of US$8.2 bn [3, 4]. The aim of this chapter is to highlight the applications of textiles in medical and applied healthcare sectors.

15.2 Textiles for biomedicine

15.2.1 Scaffolds for tissue engineering

Transplant surgery is the leading therapy to treat the patients suffering from organ failure or tissue loss. In 2001 in the United States alone, despite
24,076 lifesaving organ transplants, 6439 people died waiting for a transplant, leaving 84,798 registrations on the waiting list for an organ at the end of the year [5]. It is clear that organ transplant alone is not a viable solution to treating organ failure and tissue loss. There is a useful overview of the use of biotextile scaffolds for tissue engineering applications, demonstrating how polymer chemistry, fibre science, textile technology and engineering were integrating to make significant contributions to novel designs [6]. It was back in the 1980s that Professor D. F. Williams of the University of Liverpool, UK, first defined the term biomaterial as a nonviable material used in the fabrication of a medical device and intended to react with biological systems [7]. Following the same line of thinking, the term biotextiles was defined as a structure composed of textile fibres and designed for use in specific biological environment (e.g. surgical implants), where its performance depends on its interactions with cells and biological fluids as measured in terms of its biocompatibility and biostability [8].

Table 15.2 [9] is a partial list of some of the most common implant applications of textiles. It has been generally accepted that healing occurs more rapidly if an inert biomaterial is used which the body can tolerate more readily. Biomedical implants are used to aid or replace damaged tissues or organs. These materials are used in effecting repair to the body whether it is wound closure (sutures) or replacement surgery (vascular grafts, artificial ligaments, etc.) [10].

Tissue engineering may be defined as the applications of the principles and methods of engineering and lifesciences towards fundamental understanding of the structure function relationship in normal and pathological mammalian tissues, and the development of biological substitutes for the repair and regeneration of tissue and organ function [11, 12]. This work was pioneered by Langer and Vacanti in the early 1990s who summarized progress with matrices, cells, in vitro bioreactor systems, and the generation of devices with their own vascular supply (Figure 15.1). The strategy of tissue engineering generally involves the following steps:

(a) An appropriate cell source must be identified, isolated, and produced in sufficient numbers.

(b) An appropriate biocompatible material that can be used as a cell substrate (open system) or cell-encapsulation material (closed system) must be isolated or synthesized and manufactured into the desired shape and dimensions.

(c) The cells must be uniformly seeded onto or into the material and grown in a bioreactor.

(d) The engineered structure is placed into the appropriate in vivo site. Depending on the site and the structure, vascularization may be necessary [13].
Systems have been designed to be either open or totally integrated into the recipient, or closed (encapsulated) to provide protection from the host’s immune system. Tissue-engineered devices can also use controlled drug-delivery methods to release growth factors that may augment angiogenesis or aid in new tissue generation [14–16]. A critical step of all tissue engineering techniques is the use of a tridimensional structure which, mimicking the extracellular matrix (ECM), serves as scaffold which is able to promote and guide actively the tissue regeneration process. The ability of the scaffold in releasing signalling molecules, such as growth factors (GFs), in a

<table>
<thead>
<tr>
<th>Application</th>
<th>Materials</th>
<th>Textile structures [monofilament (m), yarn (y), woven (w), braided (b), knitted (k), and nonwoven (n)]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal wall</td>
<td>Polyester, polytetrafluoroethylene (PTFE), polyurethane</td>
<td>w, k</td>
</tr>
</tbody>
</table>
| Blood vessel (vascular graft) | Polyester  
Bone plant | Carbon, PGA  
Cartilage | Low density polyethylene, polyester, PTFE, carbon | w, b  
Dental bridge | Ultrahighmolecular weight polyethylene (UHMWPE), carbon, glass, aramid | w, n  
Dental post | Carbon, glass  
Dural substitute | Polyester, PTFE, polyurethane, collagen | y, b  
Heart valve (sewing ring) | Polyester | k, w  
Intervertebral disc | Polyester, PTFE | w, k  
Intramedullary rod | Carbon, glass | y, b  
Joint | Polyester, carbon, UHMWPE | w, k  
Ligament | Polyester, carbon, glass, aramid | b, w, k  
Orthodontic arch wire | Glass | y, b |
Skin | Chitin | n, w, k  
Spine rod | Carbon | y, b  
Suture | Polyester, PTFE, polyamide, polypropylene, polyethylene, collagen, polylactic acid (PLA), polyglycolic acid (PGA) | m, y, b  
Tendon | Polyester, PTFE, polyamide, polyethylene, silk | b, w, y
Polyester and nylon based textiles in biomedical engineering

controlled fashion is critical to achieve a successful tissue development and repair [17]. Hence, there has been a multitude of research work carried out in the last decade to design and develop various types of optimum scaffolds for tissue engineering (Table 15.3).

Scaffolds play a central role in tissue engineering [18, 19]. Textile structures are particularly attractive to tissue engineering because of their ability to tailor a broad spectrum of scaffolds with a wide range of properties. Textiles are interesting for biodegradable tissue engineering scaffolds. The cellular components will generate new tissue through production of an extracellular matrix, while the scaffold material provides structural integrity and mechanical stability during this process. Scaffold structure and porosity are key elements that will govern the formation of new tissue and subsequent neovascularization in vivo. There is a need for structural biocompatibility of the scaffold and the host tissue [18]. The optimum design of a scaffold for a specific tissue application requires consideration of microstructural, chemical and biological aspects. It is often difficult to isolate these aspects as they are interdependent and sometimes their effects are unknown.

Microstructural aspect

The microstructural aspect of scaffolds includes pore size, porosity, pore size distribution, pore connectivity and reproducibility of pores. These aspects are vital, as they provide the optimal spatial and nutritional conditions for the cells, and determine the successful integration of the natural

* Cells may be tissue specific, stem cells, or embryonic stem cells. They may be autologous or allogenic
† The matrix may be natural or synthetic, it may be fibrous, a foam, a hydrogel, or capsules
‡ In-vitro culture may be in static, stirred, or dynamic flow conditions

15.1 Tissue engineering process [14].

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Table 15.3 Various scaffolds used in tissue engineering [9]

<table>
<thead>
<tr>
<th>Tissue engineered biological substitute</th>
<th>Scaffold material</th>
<th>Scaffold structures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bladder</td>
<td>PGA</td>
<td>Textile (n)</td>
</tr>
<tr>
<td>Blood vessel</td>
<td>Polyester (Dacron), polyurethane, ePTFE, PGA, PLA, PGLA (Vicryl) Collagen</td>
<td>Textile (n, w, b, k)</td>
</tr>
<tr>
<td>Bone</td>
<td>PGA, PLGA + hydroxyapatite fibres PLLA</td>
<td>Textile (n), Foam</td>
</tr>
<tr>
<td>Cartilage</td>
<td>PGA, PLLA, PGLA</td>
<td>Foam Textile (n)</td>
</tr>
<tr>
<td>Dental</td>
<td>DL-PLA, PGLA (Vicryl)</td>
<td>Foam (porous membrane), Textile (n)</td>
</tr>
<tr>
<td>Heart valve</td>
<td>PGA</td>
<td>Textile (n, w)</td>
</tr>
<tr>
<td>Tendon</td>
<td>PGA</td>
<td>Textile (n, y)</td>
</tr>
<tr>
<td>Ligament</td>
<td>Collagen</td>
<td>Textile (y)</td>
</tr>
<tr>
<td></td>
<td>PGA, PLAGA</td>
<td>Textile (b, n)</td>
</tr>
<tr>
<td>Liver</td>
<td>PGA, PLA, PGLA, polyorthoesters, polyanhydride PLGA</td>
<td>Foam, Textile (n)</td>
</tr>
<tr>
<td>Nerve</td>
<td>Collagen-glycosaminoglycan, PGA</td>
<td>Foam, Textile (n)</td>
</tr>
<tr>
<td>Skin</td>
<td>PGA, PGLA (Vicryl), Nylon Collagen-glycosaminoglycan</td>
<td>Textile (w) Foam</td>
</tr>
</tbody>
</table>

Note: PLAGA, PGLA, PLGA are copolymers of polyglycolic acid (PGA) and polylactic acid (PLA).

tissue and the scaffold. For example, Zeltinger et al. [20] studied the influence of two key scaffold design parameters, void fraction (VF) and pore size, on the attachment, growth, and extracellular matrix deposition by several cell types on disc shaped, porous L-PLA scaffolds with two VF (75% and 90%) and four pore size distributions (<38, 38–63, 63–106, and 106–150 μm). DmFb (canine dermal fibroblasts), VSMC (vascular smooth muscle cells), and MVEC (microvascular epithelial cells) showed uniform seeding on scaffolds with 90% VF for each pore size, in contrast to the
corresponding 75% VF scaffolds. Culture data from scaffolds with a 75% VF suggests that the structural features were unsuitable for tissue formation. Hence, there were limits of acceptable scaffold architecture (VF, pore size) that modulated in vitro cellular responses. Table 15.4 compares the various microstructural aspects of foams and textile structures.

Lee et al. [21] investigated the effect of interconnectivity of pores on cell attachment and proliferation as well as surface properties of PLGA scaffold for skin tissue engineering. The interconnectivity of pores determines the transport of nutrients and waste and thus influences the success of tissue engineering [22]. The reproducibility of scaffolds is also very important as it determines the dimensional stability of the scaffold as well as the consistency of tissue formation. In a typical textile scaffold, three levels of porosity can be achieved. The arrangement of fibres in the yarn determines the accessible space for cells. The inter-fibre space (or groove between two adjacent fibres) may be considered as the first level of porosity. It has been found that the fibroblasts preferentially organize themselves along the length of the PET fibres, grouping along the groove created by two adjacent fibres. It is interesting to see that fibroblasts are capable of bridging fibres which are as far as 40 μm apart. The inter-fibre gap or first level of porosity in a textile scaffold can be controlled by changing the number of fibres in the yarn and also the yarn packing density. Further variations in porosity can be achieved by using twisted, untwisted, textured, untextured, continuous or spun yarns.

The gap or open space between the yarns (it is open space inside the loop in the case of knits) forms the second level of porosity. In the case of knitted scaffolds, the porosity can be varied selectively by changing the stitch density and the stitch pattern. In the case of braided scaffolds, porosity can be varied by controlling the bias angle of the interlacing yarns. For woven scaffolds, it is possible to change the porosity by controlling the inter-yarn gaps through a beating action. During the seeding on woven, braided and knitted scaffolds with hepatocytes, it has been observed that the cells attach preferentially at the inter-yarn gaps or pores in woven and braided scaffolds, whereas they clump together on the ridges of curved yarns in the case of knitted scaffolds.

It may be noted that woven and braided scaffolds share similar surface topographies formed by the interlacing yarns. Knitted scaffolds, however, comprise curved yarns, which have a significant effect on the behaviour of hepatocytes. Unlike hepatocytes, the fibroblasts attach to the ridges of yarns irrespective of the scaffold type. The different behaviour of fibroblasts and hepatocytes may be due to their different cell sizes and shapes. It may be noted that the diameter of fibroblasts ranges from 10 μm to 20 μm, and they flatten out after attachment. The hepatocytes are larger with diameters in the range 15 μm to 30 μm, and they retain their spherical
<table>
<thead>
<tr>
<th>Fabrication</th>
<th>Foam/sponge</th>
<th>Nonwoven</th>
<th>Woven</th>
<th>Braided</th>
<th>Knitted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pore size (μm)</td>
<td>0.5–500</td>
<td>10–1000</td>
<td>0.5–1000</td>
<td>0.5–1000</td>
<td>50–1000</td>
</tr>
<tr>
<td>Porosity (%)</td>
<td>0–90</td>
<td>40–95</td>
<td>30–90</td>
<td>30–90</td>
<td>40–95</td>
</tr>
<tr>
<td>Pore distribution</td>
<td>Random to uniform</td>
<td>Random</td>
<td>Uniform</td>
<td>Uniform</td>
<td>Uniform</td>
</tr>
<tr>
<td>Reproducibility of porosity</td>
<td>Poor to good</td>
<td>Poor</td>
<td>Excellent</td>
<td>Excellent</td>
<td>Good to excellent</td>
</tr>
<tr>
<td>Pore connectivity</td>
<td>Good</td>
<td>Good</td>
<td>Excellent</td>
<td>Excellent</td>
<td>Excellent</td>
</tr>
<tr>
<td>Processability</td>
<td>Good</td>
<td>Good</td>
<td>Excellent</td>
<td>Excellent</td>
<td>Good</td>
</tr>
<tr>
<td>Other comments</td>
<td>Current techniques are associated with processing undesirable residues such as solvents, salt particles</td>
<td>Equipment cost is high. Control over porosity is always questionable</td>
<td>Shapes are limited</td>
<td>Limited to tubular or uniform cross-sectional shapes</td>
<td>Limited by the low bending properties of current biodegradable fibres</td>
</tr>
</tbody>
</table>

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structure even after attachment to the scaffold. Furthermore, a third kind of porosity can be introduced by subjecting the textile structures to secondary operations such as crimping, folding, rolling, stacking, etc. In other words, it may be stated that the flexibility of microstructural parameters is tremendous in the case of textile scaffolds [9].

**Mechanical aspect**

The mechanical aspect of scaffolds, such as structural stability, stiffness and strength, have considerable influence on cellular activity. For example, in tissues like bone, cell shape is influenced by mechanical forces. Cell shape modification takes place as a result of external forces including gravity, and also of internal physical forces. Cell shape modification also depends on the nature (constant or cyclic), type (uniaxial, biaxial, multiaxial, etc.) and magnitude of the mechanical stimulation. Mechanical stimulation also affects the release of soluble signalling factors and the deposition of extracellular matrix constituents [9, 23]. Researchers are making use of these observations in the case of bone tissue engineering. They are applying external mechanical stimulation to promote tissue formation [22, 24]. Therefore, in bone tissue engineering, the scaffolds are designed to withstand severe physiological loads [24]. In blood vessel applications, the scaffold needs to be strong enough to withstand physiologically relevant pulsatile pressures and at the same time match the compliance or elasticity values of a native blood vessel [25, 26]. The mechanical aspects of various scaffolds are compared in Table 15.5. Of all the scaffolds, knits display considerable deformability and good compliance owing to their looped yarn arrangement. Hence, they are suitable for bladder [27, 28] and blood vessel tissue engineering applications [29].

**Biological aspect**

There is increasing evidence that scaffold surface chemistry influences cellular activity. The surfaces with amine groups are best for the CHO (Chinese hamster ovary) cell adhesion, spreading and growth probably owing to the positively chargeable character in an aqueous cell culture medium; a large portion of cell or serum protein surface is recognized as being negatively charged, resulting in electrostatic interaction between the surfaces. For surfaces with neutral functional groups; hydroxyl groups showed better cell spreading than amide groups, probably due to specific hydrogen bonding between the surface hydroxyl groups of the polymer and the polar groups of the cell surfaces. So the surface with COOH group showed poor cell adhesion due to the presence of negative charge [30]. Boyan et al. [31] showed that osteoblast response varies with the material
**Table 15.5 Mechanical aspects of the scaffolds [9]**

<table>
<thead>
<tr>
<th>Fabrication</th>
<th>Foam/sponge</th>
<th>Nonwoven</th>
<th>Woven</th>
<th>Braided</th>
<th>Knitted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stiffness</td>
<td>Low</td>
<td>Low</td>
<td>High</td>
<td>High</td>
<td>Medium</td>
</tr>
<tr>
<td>Strength</td>
<td>Low</td>
<td>Low</td>
<td>High</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>Structural stability</td>
<td>Good</td>
<td>Poor to good</td>
<td>Excellent</td>
<td>Excellent</td>
<td>Poor to good</td>
</tr>
<tr>
<td>Drapeability</td>
<td>Poor</td>
<td>Isotropic behaviour</td>
<td>Good</td>
<td>Isotropic behaviour</td>
<td>Excellent</td>
</tr>
<tr>
<td>Other comments</td>
<td>Anisotropic, with good properties parallel to fibres and poor properties normal to fibres</td>
<td>Poor Anisotropic, with good properties in axial direction and poor properties in transverse direction</td>
<td>Excellent</td>
<td>The behaviour can be tailored from isotropic to anisotropic</td>
<td></td>
</tr>
</tbody>
</table>

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on which cells are cultured, and attributed this to differences in the surface chemistry, charge density and net polarity of the charge. Van Wachem et al. [32] investigated the in vitro interaction of human endothelial cells (HEC) and polymers with different wetabilities in a culture medium containing serum. Optimal adhesion occurred onto moderately wetable polymers.

Modification of synthetic polymeric scaffolds by natural polymers (components of extracellular matrix (ECM)) like collagen [33–36], gelatin [2, 37], fibronectin [38], laminin [39] improve their biological behaviour. Since these biopolymers belong to animal origin, so they are not antigenic and immunogenic, and they have high affinity for cell adhesion and their proliferation due to presence of specific peptide sequence. So, the combination of various factors, such as scaffold material, structure, physical, chemical, mechanical, and biological properties, cell types, in vitro or in vivo conditions, etc., determines the success of tissue engineering.

Fabric structure and design aspect

Textile fibres, yarns, fabrics, composites and 3-D shaped fabrics from woven, knitted, nonwoven, braided and embroidery processes play a vital role in the manufacture of various implants, including the replacement of diseased or non-functioning blood vessels and segments of the aorta or other big arteries. It is even feasible to produce vascular prostheses as fine as 2 to 3 mm in diameter [10].

Woven

The first commercial prostheses were woven on two sets of yarn that had a high fabric count. This provided a type of graft which was rigid, tightly woven, and had a low permeability, therefore bleeding is reduced [40]. Risbud et al. [41] reported cell growth on the scaffolds consists of a woven PET fabric with well defined macropores and coated with biodegradable chitosan-collagen membrane. Both collagen and chitosan are shown as suitable substrates for hepatocyte attachment, growth and differentiation. The degradable polymer membrane could create a concentration and pressure gradient on two sides of the membrane that will vary with the diffusion and perfusion properties of the scaffold with degradation. Karamuk et al. [42] coated the woven PET fabric on one side with a thin biodegradable film (PLGA), in order to obtain a polar structure for developing the scaffold for liver cell culture. The development of a composite structure ensures the stability of the membrane during in vitro degradation, independent of the mesh size. Similarly, Dacron fabric may absorb cyclodextrin
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(CD) which has been observed to show optimum behaviour as vascular grafts [43].

Nonwoven

Nonwovens are open porous, three dimensional structures in which the cells can proliferate and be supplied with nutrient solution. This is an important advantage, as the surface, on which cells can grow, is very large in relation to the amount of biomaterial implanted. Nonwovens have fibrous characteristics similar to collagen which are not shared with non-textile materials. Spun bonded nonwovens are generally made by applying bonding agents or molten adhesives. These processing aids reduce pore size (but highly porous, 96% pre-volume), change surface properties and often render nonwoven nonbiocompatible. Staple fibre nonwovens are the fibres cut online and carded, laid down on a conveyor belt, strengthened by calendaring, and finally needled. But the major disadvantage associated with this is shrinkage after implantation in patients because the fibres cannot be fully oriented by online drawing resulting in incomplete crystallization. An advantage of the staple fibre nonwoven process is the possibility of mixing fibres with different resorption rates, such as resorbable with non-resorbable fibres. Microfibre fleeces are small tubular prosthesis. Porosity is less (70%) than spun bounded nonwovens, but pore size can be large (up to 300 μm). Solvent spun microfibre fleeces are small calibre vascular prostheses. Pore size can be extremely small (1 μm) but porosity is also low (60%) [44].

The novel nonwoven PLGA structures produced by the electrospinning process have been developed for tissue engineering applications. The electrospun structure with 500 to 800 nm diameter features a morphologic similarity to the extracellular matrix (ECM) of a natural tissue, which acts to support and guide cell growth [45]. Cells on such surfaces are sensitive to topography. Electrospinning can be altered to influence either the surface topography of the fibres themselves or the larger topography of the web of spun fibres [46]. For example, a change in the chain conformation in nylon 6 and nylon 12 due to electrospinning occurs, implying that a high stress is induced on the electrospinning jet as the fibres are being formed, and this stress alters the chain conformation of the nylon backbone [47]. Ma et al. [48] had also prepared polyethylene terephthalate nanofibre mats (PET NFM) by electrospinning for blood vessel engineering. The electrospun PET NFM was first treated in formaldehyde to yield hydroxyl groups on the surface, followed by the grafting polymerization of methacrylic acid (MAA) initiated by Ce(IV). Finally, the PMAA-grafted PET NFM was grafted with gelatin using water-soluble carbodiimide as coupling agent. The gelatin grafting method can obviously improve the spreading and proliferation of
the ECs on the PET NFM, and moreover, can preserve the EC’s phenotype. A bilayered tubular scaffold composed of a stiff and oriented PLA outside fibrous layer and a pliable and randomly oriented PCL fibrous inner layer (PLA/PCL) was fabricated for blood vessel tissue engineering by sequential multilayer electrospinning (ME). The resulting scaffolds achieved the desirable levels of pliability (elastic up to 10% stain) and proved to be capable of promoting cell growth and proliferation [49]. Nonwoven scaffolds based on the PET, PGA and PLA have been developed for tissue engineering of the anterior cruciate ligament and cartilage [50, 51].

Knitted

In this type of fabric, the yarn used is enveloped around each other in two directions, warp knit (lengthwise) and weft knit (transversely) (Figure 15.2 [18]). The difference between both is that the weft-knit fabric loosens and the warp-knit does not. This type of construction provides more pores in the centre of loops of the knitted yarn than the woven fabrics. The knitted prostheses can also elongate thereby affecting its dimensional stability. The knitted polyester vascular prosthesis has become the standard vascular

![Knitted polyester vascular graft structure](image)

**15.2 Knitted polyester vascular graft structure: (a) weft knitting; (b) warp knitting [18].**
graft for replacement of arterial vessels of 6 mm and greater. The knitted structure, by its nature, is porous, which is what is required for incorporation by tissue ingrowth from the host [2]. A major problem of arterial grafts (as with other vascular grafts) is the induction of clotting by the graft’s surface. To circumvent this problem, grafts are usually clotted with the patient’s own blood before implantation to reduce seepage through the knitted structure. Pre-clotting has been found useful in delaying or preventing thrombosis [2]. Branched hybrid vascular prostheses have also been developed on the base of type I collagen with minimal reinforcement by a knitted fabric mesh made of segmented polyester.

Embroidery

Embroidery is generally defined as the decoration of woven or knitted textile fabrics or other surfaces, e.g. leather, through the application of threads or other decorative objects (beads, cords, applications), by sewing them in or on, in an arrangement designed to achieve a pattern on the ground fabric [52]. What was until recently unthinkable has now come about: medicine has discovered embroidery. This is particularly advantageous with relatively small motifs. As opposed to weaving, where threads are arranged at rigid angles, embroidery also enables rounded patterns. Additionally, made-up embroidery goods are dimensionally stable – unlike knitted fabrics. This technology allows implants to be constructed in such a way that embroidered, three-dimensional structures become functional in tissue engineering [52]. Hernia patches, implants for intervertebral disc repair and a graft stent for the repair of aortic aneurysm have been designed by Ellies and coworkers [53].

In the development of medical textiles, polyester is frequently used. If specialized biocompatible materials are used, they are brought in at a later, more advanced stage of development, due to their high cost (up to US$4000/kg). Various synthetic or natural polymer fibres feature very specific structural and mechanical properties, favoured by tissue engineering as bone, cartilage or skin replacements [10]. Karamuk et al. [54] carried out tests with embroidered materials that decompose inside the body. These were threads made of polyglycolic acid (PGA). In vitro tests showed that forces in embroidery goods can be controlled by embroidery technology; in this way, the mechanical properties of the textile can be adapted to those of the body tissue, and inflammatory reactions can be avoided.

Braided

A braid exhibits a locking angle, i.e., an angle between the carrier bundles of the braid which when reached prevents the individual carrier bundles
from moving independently of each other thus resulting in a dramatic increase in stiffness. The scissoring effect of an opening and closing braid can damage ingrowing tissue. Braids are not the ideal structure for tissue engineering where there is expected to be regular loading and un-loading of the developing tissue [56]. Plain tubular braids may be used as prostheses for the replacement of injured ligaments in joints, like the human knee joint [57, 58]. The simple reciprocal relation of braiding position to pick counts allows an easy determination of the limits of the braiding machine on the design and manufacturing of braids. This is important for the calculation of the stress-strain behaviour of the prosthesis, which should be adapted to the individual situation within the joint as well as to the intended implantation position [57]. Irsale et al. [58] focused on prototype manufacturing of polymeric stents. The prototype is an integrated braided and tubular narrow woven fabric assembly.

The braided structure manufactured with polyester monofilaments acts as the reinforcing component, and the tubular narrow woven fabric tightly covering it acts as the sealing component. Prototypes of bifurcated braided stent for abdominal aortic aneurysm applications are also manufactured with braiding of polyester monofilaments [58].

Composite materials

Textile composites are produced by impregnating matrix materials into their dry preforms to hold the multidirectional yarns together. This is generally done by liquid moulding techniques such as resin transfer moulding, structural reaction injection moulding, and resin film infusion. For example, Peltola et al. [59] prepared a composite from P(L/DL)LA by the sol-gel method. Fibres having active surface properties or suitable porous structure can be used as such or as a bioactive part in composites. Braided fabrics can be used in textile composites e.g. bone plates [60]. The integral structures of braided textile composites enable them to endure twisting, shearing and impact better than do woven or knitted fabrics. Due to their higher impact resistance/tolerance and stability or conformability under tension in the braided yarn system, the braided fabrics can be designed for multidirectional conformity [61]. Branched hybrid vascular prostheses have been developed on the base of type I collagen with minimal reinforcement by a knitted fabric mesh made of segmented polyester. The inner diameter was prepared by pouring a cold mixed solution of bovine smooth muscle cells and collagen into a corresponding tubular mould and by subsequent thermal gelatination, followed by seven-day culturing. Reinforcement with an elastomeric mesh improved mechanical strength of the hybrid tissue and created compliance matching with native arteries. A branched or bifurcated hybrid graft with mesh reinforcement is expected to be
applicable to arterial replacement in a branching region [62]. Huang et al. [63] evaluated the mechanical properties of the multilayered knitted fabric-reinforced composite laminates. These understandings are believed to be useful in the design of composite structures. Moutos et al. [64] presented a micro-scale 3D weaving technique to generate anisotropic 3D woven structures as the basis for novel composite scaffolds that are consolidated with a chondrocyte-hydrogel mixture into cartilage tissue constructs. Other biomedical applications of textile reinforced composites are dentistry and orthopaedics [65].

Surface activation for bifunctionalization of scaffold

Textile materials offer the possibility to combine compliance and porosity in scaffolds for organ reconstruction where it can support the three dimensional growth of cells to produce functional tissue [27]. Scaffolds have also been made from extracellular protein matrix components, such as collagen, laminin or fibronectin. These materials show excellent cell adhesion, biodegradability and biocompatibility but suffer from the disadvantage that they cannot be freely or reproducibly processed into stable objects with three dimensional shapes of good mechanical strength. Conversely, scaffolds made from thermoplastic polymers have excellent strength and ductility and can be processed into various shapes and products, but do not have surfaces that readily interact with cells. These textile surfaces therefore need to be modified in such a way that the surface acquires chemical functionality that may attract extracellular matrix proteins and later cells [27]. The surface modifications of synthetic polymers are necessary to make them bioreceptive. Various researchers worked on surface modification of different types of biomaterials used for various purposes. Commonly techniques used for surface modifications are as follows.

Grafting process

Grafting technology has long been known in polymer chemistry, but the modification of the polymer surface by grafting is a fairly new technology. Two methods are available for the grafting of a polymer surface: coupling reaction of existing polymer chains; and graft polymerization of monomers on to a preexisting polymer.

Plasma induced grafting on polyester

When polymeric materials are exposed to plasma, radicals are created in the polymeric chains. These radicals can initiate reactions when they are in contact with monomers in a liquid or gaseous phase. Inelastic collisions
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of electrons in plasma with a polymer surface generate radicals at the surface of a polymer through excitation of the polymer molecules. As a result, a grafted chain is formed on the surface of the polymeric material. The grafted surfaces may provide active sites for the binding of protein molecules. For instance, the poly (acrylic acid) grafted PET surface may be immobilized with collagen by a dip coating process [27, 33]. The collagen immobilized surface provides an excellent surface for the growth of smooth muscle cells. Cells adhere on the surface and grow rapidly. In an another study, Bisson et al. [37] proposed a scaffold model for bladder reconstruction by plasma induced graft polymerization of acrylic acid on PET films which subsequently allowed collagen (Type I and Type III) immobilization and human smooth muscle cell expansion.

PAA-grafted PET films, onto which serum proteins of the culture medium adsorbed spontaneously, proved to be better matrices than films on which collagen has been immobilized. Insulin and heparin co-immobilized PET (PET-I-H) was also prepared by grafting of polyethylene oxide (PEO) onto PET-AA followed by reaction first with insulin then with heparin (Figure 15.3) [66]. Hsu et al. [67] prepared lactide grafted

15.3 Schematic diagram showing the immobilization of insulin and/or heparin on PETs [66].
polyurethanes by exposing the polyurethane films to argon plasma discharge, followed by grafting L-lactide onto the plasma treated surface. The grafted surfaces showed enhanced attachment and growth in both 3T3 fibroblast and human umbilical vein endothelial cell culture tests. L-lactide monomers grafted onto polyurethane substrates could therefore be useful in facilitating the endothelial cell seeding process in small vascular applications.

Radiation grafting on polyester

The material surface may be designed to exhibit a desired physico-chemical nature by selection of monomer and grafting procedure. While gamma radiation [68, 69] induced grafting leads to modification of both the surface and the bulk, UV, ozone, and plasma modification only affect the surface leaving the bulk intact. PET surfaces prepared by grafting using these techniques have for instance been evaluated as biomaterials for artificial hair transplants [70] and for the immobilization of biomolecules such as heparin and insulin [66]. Knitted PET fabric has been modified by radiation induced graft polymerization of methacrylic acid (MAAc)/N-vinylpyrrolidone (NVP) monomer. The influence of reaction conditions on the degree of grafting and the physical structure of the grafted fabric was investigated in this study. The grafted surfaces have also been evaluated for collagen immobilization and for seeding of urothelial cells with the ultimate aim of using this for applications in urology. Jou et al. [71] also grafted acrylic acid (AA) on PET fibre by a radiation method. The resulting fibres were further grafted with chitosan (CS) via esterification followed by collagen (COL) by gluteraldehyde (GA). The results indicate that by grafting with CS and immobilizing with COL, PET fibres exhibited both antibacterial activity against pathological bacteria and improvement in the proliferation of fibroblast.

15.2.2 Sutures

Sutures are probably the largest group of devices implanted in humans and can be used in skin, muscle, fat, organs and vessels. Although they seem to be of small concern to the medical community, few devices have been made of so many different materials. The suture market currently exceeds US$1.3 bn annually. By definition: a suture is a thread that either approximates and maintains tissues until the natural healing process has provided a sufficient level of wound strength or compresses blood vessels in order to stop bleeding [72]. The United States Pharmacopoeia (USP), European Pharmacopoeia (EP) and British Pharmacopoeia (BP) are the official compendia for the suture industry, which sets standards and guidelines for
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Properties of sutures

The ideal suture is strong, handles easily and forms secure knots. It is important that the suture causes minimal tissue inflammation and does not promote infection. It should be able to stretch, accommodate wound oedema and recoil to its original length with wound contraction. It should also be inexpensive. Since no single suture possesses all of these features, it is the physician’s task to weigh the advantages and disadvantages of the available suture materials [74]. The comparative information on the performance of various sutures is summarized in Table 15.7.

Knot strength

It is a measure of the amount of force necessary to cause a knot to slip and is directly related to the coefficient of friction of a given material [76].

Table 15.6 Suture sizes

<table>
<thead>
<tr>
<th>P</th>
<th>Diameter</th>
<th>Tensile strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>4-0</td>
<td>0.2 mm</td>
<td>7.5 N</td>
</tr>
<tr>
<td>3-0</td>
<td>0.3 mm</td>
<td>12.3 N</td>
</tr>
<tr>
<td>2-0</td>
<td>0.35 mm</td>
<td>19.6 N</td>
</tr>
<tr>
<td>0</td>
<td>0.4 mm</td>
<td>22.3 N</td>
</tr>
<tr>
<td>1</td>
<td>0.5 mm</td>
<td>37.3 N</td>
</tr>
</tbody>
</table>

Table 15.7 Performance comparison of various suture materials [75]

<table>
<thead>
<tr>
<th>Sutures</th>
<th>Tensile strength</th>
<th>Tissue reactivity</th>
<th>Handling</th>
<th>Knot security</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nylon, monofilament</td>
<td>High</td>
<td>Low</td>
<td>Poor</td>
<td>Poor</td>
<td>High</td>
</tr>
<tr>
<td>Nylon, braided</td>
<td>High</td>
<td>Moderate</td>
<td>Good</td>
<td>Fair</td>
<td>Fair</td>
</tr>
<tr>
<td>Polyester</td>
<td>High</td>
<td>Moderate</td>
<td>Good</td>
<td>Good</td>
<td>Fair</td>
</tr>
<tr>
<td>Polyglycolic acid</td>
<td>Good</td>
<td>Low</td>
<td>Fair</td>
<td>Good</td>
<td>Low</td>
</tr>
<tr>
<td>Polyglycolide-lactide</td>
<td>Good</td>
<td>Low</td>
<td>Good</td>
<td>Fair</td>
<td>Low</td>
</tr>
<tr>
<td>Polycaprolactone</td>
<td>Good</td>
<td>Low</td>
<td>Poor</td>
<td>Poor</td>
<td>High</td>
</tr>
</tbody>
</table>
Bayraktar et al. [77] investigated the knot performance of various sutures and observed that knot performance depends on the surface properties. For a braided structure, the coefficient of friction is higher because the threads in the braids have mobility, which increases the knot holding capacity. Therefore, there is no knot untying before the break for the braided structures. Monofilament sutures however have very smooth surfaces, which allow them to pass easily through the tissue. At the same time, it decreases the knot holding capacity.

Elasticity
This refers to the intrinsic tension generated in a material after stretching, which causes it to return to its original length. Elasticity is a desirable feature, since it allows the suture to expand during wound oedema without causing strangulation or cutting of tissue, and to recoil during wound retraction, thereby maintaining wound edge apposition [74].

Memory
This refers to the inherent tendency of a suture material to return to its original shape after being manipulated and is a reflection of its stiffness. A suture with a high degree of memory is stiffer, more difficult to handle and more likely to become untied compared with suture material that has less memory.

Tissue reactivity
This refers to the inflammatory response generated by the presence of suture material in the wound. This response peaks within two to seven days and is a function of the quantity of material present as well as its type and configuration [76, 78, 79] Everett’s [80] studies have shown that an excessive inflammatory reaction may lead to the softening of surrounding tissues and result in decrease in wound strength. It was also observed that nylon and polyester sutures produce the strongest inflammatory reaction during the first five days. Sutures of superior tensile strength and knot security not only minimize the risk of suture line disruption but also reduce the amount of foreign material left in the wound by allowing the use of finer sutures and fewer knots. This, in turn, reduces tissue reaction and infectious complications [74]. Because of the various features, the choice of suture material for any given wound closure should not be made arbitrarily, but rather with careful attention to the physical and handling properties of the suture, as well as its propensity for eliciting tissue reaction and promoting infection [74].
Classification of sutures

Sutures can also be classified into two categories: absorbable and non-absorbable. The application of the two categories depends on the type of procedure and the physician’s preference. Table 15.8 shows the absorbable and non-absorbable sutures [81].

Absorbable sutures

Absorbable sutures have the ability to be ‘absorbed’ or decomposed by the natural reaction of the body to foreign substances. They are used internally and are designed to lose strength gradually over time by chemical reactions such as hydrolysis. It is important to note that not all absorbable sutures have the same resistance level to absorption, but each can be formulated or treated in order to obtain a desired decomposition rate and be excreted in urine or faeces, or carbon dioxide in expired air [82–84]. Currently, the most commonly used absorbable sutures are synthetic substances: polyglycolic acid and polyglactic acid.

Nonabsorbable sutures

Nonabsorbable sutures are those, which do not lose their tensile strength for a long period of time. Generally, these sutures are used for closing cutaneous or oral incisions where the sutures can be easily removed e.g. nylon, braided polyester.

Polyglycolic acid suture

Introduced in 1970, polyglycolic acid (Dexon) (Figure 15.4), a polymer of glycolic acid, was the first synthetic absorbable suture. It was hailed for its excellent tensile strength and knot strength, as well as delayed absorption and markedly diminished tissue reactivity compared with catgut. In animal studies, the absorption of polyglycolic acid suture was found to be about 40% after seven days [85]. By 15 days, it has lost more than 80% of its original strength [86, 87]. By 28 days, this material retains only 5% of its original tensile strength, and it is completely dissolved by 90 to 120 days [88]. Polyglycolic acid is absorbed by hydrolysis, reducing the inflammatory response. As a monofilament, Dexon is stiff and difficult to work with. Therefore, it is available in braided form for easier handling. Dexon also

\[
\left\{\text{OC—CH}_2—\text{O}\right\}_n
\]

15.4 Polyglycolic acid.
Polyesters and polyamides

- Polyesters and polyamides come with a synthetic coating (Dexon Plus) to facilitate known typing and passage through tissue.

Polyglactic acid suture

Polyglactic acid (Vicryl) (Figure 15.5), the second synthetic suture material to be marketed (in 1974), is a copolymer of lactide and glycolide, manufactured with a coating composed of polyglactin 370 and calcium stearate. This lubricant coating gives Vicryl excellent handling and smooth typing properties. Technical studies have shown Dexon to have slightly greater tensile and knot strength than Vicryl [89], but the differences are clinically insignificant. Like Dexon, Vicryl material retains only 8% of its original tensile strength by 28 days. However, complete absorption of Vicryl is more rapid, occurring between 60 and 90 days [87]. Like all synthetic polyesters, Vicryl degrades by hydrolysis and causes minimal tissue reaction. Vicryl is a braided suture, and comes in violet-dyed and undyed forms. When used in skin surgery, the dyed form can sometimes be seen beneath the skin surface. A buried Vicryl or Dexon suture may occasionally be extruded through the suture line.

Polyamides

Polycapramide (nylon 6) and polyhexamethylene adipamide (nylon 66) are the preferred polyamides. The other homopolymers, such as polyhexamethylene sebacamide (nylon 610), polydodecanamide (nylon 12) and polyhexamethylene isophthalamide (nylon 61), and copolymers and blends can be used. The polyamides used to make the sutures have a high molecular weight of 48000–55000. The tensile strength of the sutures varies between 132 KPSI to 145 KPSI. The sutures can be used as monofilaments or made into braided multifilament [90–93]. As a monofilament, nylon (Ethilon, Dermalon) is the most widely used non-absorbable suture in skin surgery. It has high tensile strength, minimal tissue reactivity, excellent elastic properties and low cost. The major drawback to nylon is its high degree of memory. A greater number of knot throws (three or four) are required to hold a given stitch in place.

Nylon hydrolyzes at a slow rate. Studies in rabbits have shown that buried nylon retains 89% of its tensile strength at one year and 72% by two years.
At this point, degradation apparently stabilizes; Nylon sutures retain approximately two-thirds their original strength after 11 years [95]. Thus, nylon should not be classified as a true nonabsorbable suture but rather as a very slowly absorbable suture.

Polyester suture

Braided sutures are generally obtained by using a circular braiding machine. The number of yarns composing the braid depends on the required diameter of the suture [96]. Polyester yarns were made from filaments having a weight-average molecular weight of more than 35000, a tenacity of about 7–8.5 g/denier, an elongation to break of less than 35% and shrinkage of about 0.5–2% in boiling water. The filament should have shrinkage in hot air at 3–5% (compared with its original length). The commercially available filaments meeting these requirements include Trevira High Tenacity type 712 and 787 polyester yarns from Hoechst. The polyester yarns are made up of such filaments having a denier preferably in the range 1.4–3.1. The polyester filaments are extruded in bundles (yarns) having a denier preferably of 20–350. The yarns are twisted and then braided into sutures using conventional constructions that have a sheath and, optionally, a core. However, in order to provide the best combination of suture properties, the sutures may be braided using the preferred constructions shown in Table 15.8.

The braided suture can be hot-stretched at a temperature of 160–250°C to 9–28% of its original length. The suture may also be coated with a material such as 0.1–10 wt% polybutylate, which is applied using a suspension drop coating system [97]. Braided polyester sutures are often used in cardiovascular surgery where a strong, non-absorbable suture is needed to help permanently repair tissue. However, surgeons have complained of suture roughness, which can sometimes make it difficult to secure a knot. The application of a lubricious coating can reduce suture roughness, but

| Table 15.8 Preferred constructions for a braided polyester suture [97, 98] |
|------------------|------------------|------------------|------------------|------------------|
| **Suture size** | **Sheath yarns** | **Carriers** | **Core denier** | **Picks** |
| 2                | 90–150           | 12–18          | 380–500         | 40–50          |
| 1                | 55–90            | 14–18          | 310–360         | 40–50          |
| 0                | 60–80            | 14–18          | 180–250         | 40–50          |
| 2/0              | 45–65            | 14–18          | 80–140          | 40–50          |
| 3/0              | 35–55            | 10–14          | 40–50           | 40–50          |
| 4/0              | 35–55            | 6–10           | –               | 40–50          |

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too much coating can make the suture slippery to handle and can result in tied knots coming undone.

**Developments in sutures**

The concept of utilizing a suture as a drug delivery system has been a subject of great interest in modern surgery. The delivery of antimicrobial agents near the wound closure coupled with slow release ability results in a remarkable improvement in the healing process. Sutures themselves have very little or no intrinsic biological activity, but when associated with antimicrobial agents, the entire complex molecule behaves as a germicide reservoir to prevent proliferation of a microorganism. A variety of antimicrobial agents are known which can inhibit the growth and metabolism of microorganisms. These antimicrobial agents can be bonded to the polymeric backbone either reversibly or irreversibly and are subsequently released to the surrounding medium and inhibit the growth of microbes.

**Antimicrobial sutures**

Microbes are the tiniest creatures not seen by the naked eye. They include a variety of microorganisms like bacteria, fungi, algae and viruses. In the majority of polymeric implants and intravenous catheters, Staphylococci play a predominant role. Staphylococci is the most frequently implicated microorganism in infection occurring in the drainage system of cerebrospinal fluid [99, 100], in venous catheters and sutures [101], in continuous ambulatory catheters [102], in heart valves [103] and in hip as well as knee prostheses [104]. Negative effect on the vitality of the microorganisms is generally referred to as antimicrobial. Antimicrobial agents are the chemical compounds that possess the ability to kill or inhibit the growth and metabolism of microbes thus preventing proliferation and subsequent infection [105, 106].

These can be broadly classified into two classes:

**Class I:** Those useful in treatment of disease e.g. antibiotics. Antibiotics are defined as specific chemical compounds formed by living organisms that are capable of inhibiting life processes of other organisms when used in small quantity (Table 15.9).

**Class II:** Those useful in prevention and control of diseases e.g. disinfectant and antiseptic. By convention, chemical agents, which are used to destroy microbes on inanimate objects, are disinfectants, while those applied on living tissues, especially wounds, are antiseptics (Table 15.10).
Preparation of antimicrobial sutures

The antimicrobial activity in sutures can be achieved by the following methods:

1. Blending or incorporation of volatile or nonvolatile antimicrobial agent while processing
2. Coating or absorption of the antimicrobial agent onto the filament
3. Graft polymerization followed by immobilization of antimicrobial agents onto the grafted surface.

Incorporation of bioactive agents including antimicrobial agents into polymers by blending has been commercially applied in surgical implants and other biomedical devices. This is a pretreatment technology, where the antimicrobial agent is introduced during the processing stages. For this, the additive characteristics have to be compatible with blending conditions, e.g. particle diameter, heat stability, chemical stability and effect on fibre quality [107].

Heavy metals, such as gold, silver and copper have also been used to introduce antimicrobial activity in suture [108–116]. Allard and Song [117] described another type of system in which polymer and active agent are mixed and melt together and then melt spun to a diameter greater than 1 mm. Different polymer matrix being used are polyethylene, polyester, polyglycolic acid and polyactic acid. Yabushita et al. [118] patented surgical suture with strong tensile strength. High density polyethylene and chlorohexidine hydrochloride were mixed, extruded and drawn to give monofilament with antimicrobial activity against Staphylococcus aureus (S. aureus). Of all antimicrobials, silver substituted zeolites are the most widely used to impart antimicrobial activity in polymers [119–120]. Zeolites are mainly composed of an aluminosilicate framework of alkali earth metal. Framework contains regular cavities occupied by Na+, K+, Ca2+, Mg2+, etc. These cations are easily exchangeable by other metals like Ag+, Cu2+ and Zn2+ to impart antimicrobial activity against general bacteria. These substituted zeolites are incorporated into polyesters at the level of 1–3% [121]. These metal ions inhibit the multiplication of microorganisms by two mechanisms [122]. It destroys or passes through the cell membranes and makes a bond with thiol group of cellular enzymes. As a result, it leads to alteration of the microbial metabolism and suppression of growth of microorganism. In second possible mechanism, the formation of active oxygen occurs, where these metal ions catalysed the formation of oxygen radicals that destroys the molecular structure of bacteria.

PVP-Iodine (PVP-I) complex is also one of the most widely used products in the sphere of surgery. The slow release of free iodine from the complex gives prolonged antimicrobial activity from the doped material. Antimicrobial nylon sutures have been prepared by blending of nylon 6 with PVP-I complex using a melt spinning process [123]. Depending upon the blend ratio, a pre-weighed quantity of chips was added to the saturated solution of the PVP-I complex and these chips were then spun and drawn at the temperature to obtain a monofilament with good strength. The
sutures have polar bonding between nylon and PVP-I complex, which results in the slow and controlled release of iodine from the sutures. Alternatively, silver may be used as an effective antimicrobial agent, e.g. a blend of nylon fibre and silver coated nylon fibre (known as X-static) may be used for suture application [113]. This fibre exhibits bactericidal property against *E. coli*, pseudomonas, klebsiella, staphylococcus and streptococcus species. The rate of killing (decrease of survival) increased with increase in silver ion concentration of the fibre extract, as determined through atomic absorption spectrometry.

In the post-processing technology, the most common technique for applying the antibacterial agent is coating. Antimicrobial agents that cannot tolerate the temperature used in polymer processing are often coated onto the material after fabrication. The antimicrobial agents are linked to the surface through physical bonds or anchored by the cross-linking on the fibre. Antimicrobial sutures having long lasting antimicrobial properties and good physical properties are prepared by coating a multifilament suture with a solution of an antimicrobial agent and segmented urethane polymer. Stephenson [124] coated polyester sutures with copolymer of polyquaternary polyurethane and polyanionic polymer such as heparin followed by treatment with antimicrobial agent streptomycin sulphate. The resultant antimicrobial suture gave a zone of inhibition of 0.55 cm against *Bacillus subtilis*. Antimicrobial coating on polyester suture was carried out by depositing an antimicrobial biocompatible metal by vapour deposition technique [125]. In one of the studies, it was found that immersing catgut, Dacron, silk, or chromic sutures for 24 h in a 5% or 50% aqueous solution of silver nitrate did not appreciably reduce adherence of *S. aureus*, as compared with that of unsoaked sutures [126]. Dacron and silk suture coated with a silver-zinc-allantoin complex did, however, reduce the number of adherent *S. aureus* colonies by 88% and 99%, respectively. The investigators attributed the differences in *in vitro* efficacies of the differently coated sutures to the fact that silver nitrate firmly binds to the suture material, whereas the silver-zinc-allantoin complex provides slow release of silver ions sufficient to inhibit bacterial adherence [127].

Blacker et al. [128] used silver doped bioactive glass powder (AgBr) to coat absorbable Vicryl and non-absorbable Mersilk surgical suture. Stable homogeneous coating on the surface of the suture was achieved by using an optimized aqueous slurry dipping technique. The *in vitro* bioactivity of the suture was tested by immersion in simulated body fluid (SBF). After three days of immersion in SBF, bone-like hydroxyapatite formed on the coated suture indicating the enhanced bioactive behaviour. *In vitro* antimicrobial evaluation of polyglactin 910 suture coated with triclosan has shown attractive results. The antibacterial activity of the coated suture was
evaluated against *S. aureus* and *S. epidermis* and produced zones of inhibition after five and ten passes through fascia and subcutaneous tissue. Knotted suture with triclosan gives a bacteria-free zone having volume of 14.5 cm³ for *S. epidermis* and 17.8 cm³ for *S. aureus* [129].

Iodine can be attached to polyamide sutures to provide them antimicrobial property. Antimicrobial nylon suture has been prepared by coating the monofilament with iodine [130]. Nylon 6 monofilament was treated with iodine by immersing the filament in a saturated solution of iodine in acetone and was tested for iodine release and antimicrobial properties against *E. coli* and *S. aureus* by zone of inhibition method. It was found that it showed a comparatively slower release of iodine as compared to blended suture since in coated suture iodine is present mainly on the surface so initial release is higher, then it decreases. It was also observed that a clear zone of inhibition was formed around 6 mm in the case of *E. coli* and 8 mm in the case of *S. aureus*.

Multifilament nylon suture has been made antibacterial by doping with iodine [131]. Iodine doped sutures exhibited good antibacterial activity against *E. coli*, *S. aureus*, *P. aeruginosa* and *K. pneumomoea*. The surface of the polyamide sutures has also been modified by binding of drug doxycycline [132]. The modified suture showed the release of drug into and around the wound over 10–18 days. Lin et al. [133] reported that nylon 66 fibres could be rendered antimicrobial by chemical binding of heterocyclic N-halamine functional groups to the nylon 66 molecules at the amide nitrogen, using formaldehyde as a linking agent followed by chlorination. Biocidal swatch tests showed that the nylon fabrics containing N-chlorinated hydantoin functional groups provided a 7.2 log reduction of *S. aureus* and 7.1 log reduction of *E. coli* at contact time of only 10 min, whereas unchlorinated fibres gave no reduction of bacteria even at a contact time of 71 s.

The radiation, plasma and chemical methods of grafting have occupied the attention of numerous researchers for many years [134–136]. Graft polymerization offers an effective approach to introduce desirable properties into the polymers without affecting the architecture of the polymer backbone [137, 138]. Grafting of hydrophilic monomers provides a good platform for the introduction of antimicrobial activity. The grafted side chains contain functional groups to which various bioactive materials can be attached. These functional groups include amine, carboxylic acid and hydroxyl groups, which can be utilized further for the attachment on antimicrobial drug [139, 140]. This approach of graft functionalization has been successfully used for the development of polypropylene sutures by incorporating acrylonitrile and vinylimidazole grafts into the suture matrix [141–145]. Both tetracycline hydrochloride and ciprofloxacin have been immobilized onto the modified sutures and their effectiveness against dif-
Polyester and nylon based textiles in biomedical engineering 469

different microbes has been evaluated. An antimicrobial polyester suture is prepared by graft modification of the filament in such a way that the filament retains its physical properties and attains its antimicrobial nature. In a recent study [146] the antimicrobial polyester suture was prepared by graft polymerization of acrylic acid on the surface of the suture; in order to keep the bulk unaltered grafting was carried out by using vacuum plasma. As a result, the polyester surface acquires functionality of carboxyl group while the bulk remains unaltered. It has been observed that the monomer concentration plays a crucial role in graft management and a maximum in grafting is achieved at a monomer concentration of 40% (Figure 15.6). The acrylic acid grafted surface offers excellent functionality for the interaction with chitosan molecules which is an attractive route to produce antimicrobial suture material. The chitosan coated sutures were prepared by graft modification of the filament in such a way that the filament is loaded with ciprofloxacin as the antimicrobial drug. This loading helps in the enhancement of the antimicrobial nature of the suture and hence provides better healing process. Ciprofloxacin has been observed to be released slowly from the suture and continued up to 3–4 days (Figure 15.7). This is interesting from the point of view of drug availability at the wound site till the suture is removed in 4–5 days.

Antimicrobial nylon sutures have also been developed by the graft polymerization of acrylic acid on a nylon 6 suture followed by immobilization of the antibiotics [147]. Penicillin (Pe), neomycin (Ne) or gentamycin (Ge) have been used to obtain antibacterial properties against gram-positive and gram-negative bacteria by zone of inhibition method. The release of antibiotic from the grafted suture proceeds in two stages: (I) rapid release lasts up to 1 day, which is due to absorption of drug onto the surface; (II) sustained release, lasts up to 12 days, which is due to

![Variation of degree of grafting with monomer concentration](146)
Polyesters and polyamides

Acrylic acid followed by attachment of antibiotics penicillin, neomycin or gentamycin has also been developed. It was found that the release of penicillin from PET fibre was much faster than other biocides [148].

**Barbed sutures**

In 1992, Dr Gregory Ruff of Duke University Medical Center started working on an idea of a barbed suture for cosmetic applications. Dr Ruff took the idea of a barbed suture and applied it to an absorbable suture material made of polydioxanone. The advantage of using an absorbable polymer suture is that it does not need to be removed and it does not require knots to make it secure. The knotless design has significant potential in reducing scar tissue due to the absence of a significant foreign-body reaction caused by knots. The barbed configuration anchors the suture into the tissue and provides adequate tissue adhesion while the wound heals under minimum residual tension and pressure [149]. The success of this novel wound closure device requires the suture geometry to be well characterized and monitored during manufacture for two reasons: quality control (measuring uniformity of the barb geometry) and the need to determine the effect of tissue holding capacity and the barb geometry. Quill Medical, Inc. currently produces this barbed monofilament suture from polydioxanone in size 0 (size 0 has a diameter of 0.30 to 0.39 nun), while other sizes are under development.

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The monofilament sutures contain up to 78 barbs manufactured in a spiral pattern around the circumference of the suture. The barbs are divided into two groups facing each other in opposing directions around the mid-point (Figure 15.8). The two sets of barbs divide the suture into two sections, right and left. Internal wound closure yields the best results when using an absorbable material. Using nonabsorbable wound closure material requires removal of the device after the wound has healed. This can lead to additional visits to the physician or surgeon and the use of more invasive surgical procedures. The fact that polydioxanone is absorbable makes it an ideal candidate for internal wound closure. Current sutures require the tying of surgical knots. The throws of these knots are often pushed through a transdermal cannula, which can be tedious and difficult for the surgeon as well as resulting in inferior knot performance. The knotless barbed suture can also be applied through a cannula and, without the need for tedious knot throwing and pushing, it is likely to reduce surgery time and create a more consistent method for tissue approximation.

McKenzie’s [150] article describes the use of a nylon-barbed suture for repair of the long flexor tendon of the hand (Figure 15.9). The knotless designed suture increases the flexibility and longitudinal movement of the tendon that would normally be limited by the presence of knots. The polymer suture can be engineered to maintain the required strength for the duration of the complete healing process. In addition, the absorbable barbed suture does not require removal after the repair is healed, thus reducing the number of visits to the surgeon as well as the trauma associated with a follow-up intervention.
15.2.3 Wound dressings

Wound management has recently become more complex because of new insights into wound healing and the increasing need to manage complex wounds outside hospital. The wound is a synthetic environment in which numerous cellular processes are interlinked in the process of repair. Modern dressings are designed to facilitate the function of the wound rather than just to cover it. Principles of wound dressings are changing, especially in relation to debridement of wounds and control of the wound environment. Occlusive dressings which allow debridement in a fluid environment are equal to traditional wet or dry dressings. There are some advantages in allowing wounds to heal in a moist environment which facilitates cellular migration and epithelialization. Straightforward surgical and traumatic wounds require low cost and low technology dressings. The place of more costly and complicated wound dressings has to be defined in terms of cost, labour saving and patient comfort, in addition to any putative advantages in the speed of wound healing [151]. Several studies have been carried out to develop wound dressings based on different textile fabrics. However, studies involving nylon and polyesters are very limited.

Smith and Nephew is a world leader in advanced wound care management. They are selling this wound dressing under the brand name Acticoat®. The wound dressing contains silver nanocrystals that kill bacteria in vitro in as little as 30 minutes two to five times faster than other forms of silver. The silver nanocrystal wound dressing is gaining rapid worldwide acceptance in burn and wound care centres. Its sale is continuously growing.
at double digit rates. The revenue from sales has increased from £1.7 million in 1999 to £6 million in 2003. The product is now being sold in more than 30 countries [152].

Phaneuf et al. [153] have developed a novel lightweight bioactive compression wound dressing based on polyester that provides durable infection resistance and localized haemostatic properties. Polyester (Dacron) material with polyurethane inlaid into the structure is utilized as the base material to provide the required physical properties (i.e. elasticity, durability). The surface is modified to develop functional groups. An antibiotic (ciprofloxacin) is incorporated into the material via textile dyeing technology, and a biologically active protein, thrombin, a pivotal enzyme in the blood coagulation cascade, is covalently attached to the modified surface.

15.2.4 Bandages

It should be stressed that one of the high-tech areas of medical textiles is the application of bandages for enhancing the quality of life. In fact, bandaging techniques emanate from ancient practice that facilitates further developments in improving the comfort and performance properties of value added products. Bandages are designed to perform a whole variety of special functions depending upon the type of wound and medical requirements. They can be woven, nonwoven or knitted and are either elastic or non-elastic [1].

Types of bandages

Bandages can be classified in a number of different ways but possibly the most useful method is one based upon their function as follows. The ability of a bandage to perform one or more of these functions is largely determined by its elastic properties, although the thickness, weight and conformability of the fabric are also important. Table 15.11 shows the different types of bandages, their fibre types and fabric structure [1].

Compression bandages

Compression bandages are mainly employed for the treatment of venous leg ulcers and varicose veins. It may be noted that, in the United Kingdom, there are some 400000 leg ulcer sufferers, 70% of whom have ulcers that are venous in origin [154]. The aim of the use of compression bandaging is the reduction of venous hypertension which results from valvular insufficiency [155]. The application of external compression by means of a bandage serves to increase the velocity of the blood flow within the veins.
by providing support to the muscles. It has been demonstrated that venous return is faster and more efficient if the compression bandage is applied in a manner that gives sub-bandage pressure that graduates from the ankle to the knee [156]. Elastomeric bandages made with rubber were first used in the late nineteenth century and this has now been replaced by lycra or elastane which are light, strong, comfortable and washable. These are either woven or knitted and are designed to provide prescribed levels of compression stipulated by performance based standards [157]. Compression bandages are mainly classified as elastic and nonelastic. Elastic compression bandages are categorized according to the level of pressure generated on the ankle of an average leg [158].

Light compression bandages are able to provide and maintain low levels of pressure, between 14 and 17 mmHg on an ankle of average dimensions. The clinical indications for products of this type include the management of superficial or early varices, and varicosis formed during pregnancy. In general, they are not suitable for controlling or reducing existing oedema, or for applying even low levels of pressure to very large limbs. Examples include K-Plus – Parema, and Tensolastic. Moderate compression bandages may be used to apply compression between 18 and 24 mmHg on an ankle of average dimensions. They are indicated for the treatment of varicosis during pregnancy, varices of medium severity, the prevention and treatment of ulcers and the control of mild oedema. High compression bandages may be used to apply high levels of compression between 25 and 35 mm Hg on an ankle of average dimensions. Indications for these bandages include the treatment of gross varices, post-thrombotic venous insufficiency, and the management of leg ulcers and gross oedema in limbs of average circumference. Products in this category are not necessarily able to achieve these levels of pressure on very large limbs that have been further enlarged.
by the presence of oedema. Examples include Tensopress, Setopress and Surepress. Extra-high performance compression bandages are capable of applying pressures up to 60 mmHg. The power in the bandages is such that they can be expected to apply and sustain these pressures on even the largest and most oedematous limbs for extended periods. This group includes Elastic Web Bandage BP (Blue Line Webbing) and Varico Bandage [159].

Compression can be exerted to the leg either by a single layer bandage or multilayer bandages. In the four layer system the first layer is a nonwoven padding bandage that absorbs exudate and protects bony prominences from excessive pressure. The second layer is a crepe bandage which adds absorbency and smoothes the padding layer. The third layer is a light compression bandage that is highly conformable to accommodate a difficult limb shape. The fourth layer is a cohesive flexible bandage. It applies pressure and is cohesive in nature, which means the bandages stay in place and maintain effective levels of compression for up to one week.

Studies conducted to assess the relative effectiveness of different compression bandages and compression stockings in the treatment of venous leg ulcers incorporated 27 products manufactured by different companies. The studies included comparisons between compression and no compression; between elastic multilayer high-compression bandages and non-elastic high compression bandages, as well as non-elastic multilayer high-compression bandages; between multilayer high-compression bandages and single-layer high-compression bandages; between alternative multilayer systems; and between compression stockings and compression bandages. It was concluded that compression increases ulcer-healing rates compared with the situation when there is no compression. High compression is more effective than low compression but should only be used in the absence of arterial disease. It is generally accepted that composite bandage systems are most effective when a long-stretch or short-stretch bandage system is used. A comparative study after 15 weeks' treatment of patients with venous ulceration indicated that the ulcer-healing rate for the long-stretch bandage was 58% as compared with 38% for the short-stretch bandage. Similarly, a calcium alginate dressing combined with a four-layer hydrofibre dressing group demonstrated a significantly lower frequency of dressing changes than that with alginate-fibre dressings. More patients achieved a seven-day wear time with the hydrofibre dressing than with alginate-fibre [1].

In another study, it was found that non-occlusive hydrofibre dressings were suitable for the treatment of exudating venous leg ulcers. The hydrofibre-dressing group demonstrated a significantly lower frequency of dressing changes than that with alginate-fibre dressings. More patients achieved a seven-day wear time with the hydrofibre dressing than with alginate-fibre [1].
Orthopaedic bandages

Orthopaedic bandages play a significant role in the successful treatment of venous leg ulcers. A padding of at least 2.5 cm thickness is placed between the limb and the compression bandage to distribute the pressure evenly at the ankle as well as the calf region. Wadding helps to protect the vulnerable areas of the leg from the high compression levels required along the rest of the leg [160]. Padding can also be used to reshape legs that are not narrower at the ankle than the calf. It makes the limb more like a cone-shape so that the pressure is distributed over a pressure gradient with more pressure at the foot and less at the leg. Generally the longer a compression bandage system is to remain in place, the greater is the amount of padding needed [161].

An ideal orthopaedic bandage should meet the following requirements:

- Light weight
- Soft and impart cushioning effect to the limb
- Capable of preventing tissue damage
- Good absorption and wicking properties
- Should tear easily by hand
- Comfortable and should not produce irritation or any allergic reaction to the skin on prolonged contact.

Commercial padding bandages

The most commonly used commercial padding nonwoven bandages, the fibre types and structures of the bandages are given in Table 15.12.

<table>
<thead>
<tr>
<th>Bandage code</th>
<th>Fibre type</th>
<th>Blend</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>PB1</td>
<td>Polyester</td>
<td>100%</td>
<td>Needle punched (one side)</td>
</tr>
<tr>
<td>PB 2</td>
<td>Polyester</td>
<td>100%</td>
<td>Needle punched &amp; thermal bonded</td>
</tr>
<tr>
<td>PB3</td>
<td>Viscose</td>
<td>100%</td>
<td>Needle punched (one side)</td>
</tr>
<tr>
<td>PB 4</td>
<td>Polyester/viscose</td>
<td>40%/60%</td>
<td>Needle punched (one side)</td>
</tr>
<tr>
<td>PB 5</td>
<td>Polyester/polyolefin</td>
<td>85%/15%</td>
<td>Needle punched (one side) &amp; thermal bonded</td>
</tr>
<tr>
<td>PB 6</td>
<td>Polyester</td>
<td>100%</td>
<td>Needle punched (one side) &amp; thermal bonded</td>
</tr>
<tr>
<td>PB 7</td>
<td>Polyester/viscose</td>
<td>50%/50%</td>
<td>Needle punched (one side)</td>
</tr>
<tr>
<td>PB8</td>
<td>Polyester</td>
<td>100%</td>
<td>Needle punched (one side)</td>
</tr>
<tr>
<td>PB9</td>
<td>Polyester</td>
<td>100%</td>
<td>Needle punched (one side)</td>
</tr>
<tr>
<td>PB10</td>
<td>Polyester</td>
<td>100%</td>
<td>Needle punched (one side) &amp; thermal bonded</td>
</tr>
</tbody>
</table>

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Test methods

- Bulk density (calculated from area density and thickness)
- Tear resistance
- Demand absorption and wicking (using an instrument developed at Bolton Institute).
- British Pharmacopoeia absorption method
- Sinking time
- Pressure transference (using an instrument developed at Bolton Institute).

The significance of the tests that determine the performance and properties of padding bandages is given in Table 15.13.

Effect of bulk density

Bulk density determines the bulkiness of bandages. It is important to mention that an appropriate bulkiness, say 0.05 g cm$^{-3}$, would be required to protect the bony prominences in the leg. Since padding bandage is applied next to the skin around the leg, it must be capable of imparting comfort and cushioning effect to the patient. It will be observed from Table 15.14 that the bulk density of all the commercial padding bandages are within the acceptable limit and PB1, PB3, PB4, PB7 and PB9 registered higher bulk densities.
Polyesters and polyamides

Influence of tear strength

The tear strength has a significant influence as far as the wrapping of padding is concerned. An ideal bandage is generally torn by hand after wrapping around the leg and this gives more flexibility to nurses without looking for scissors. The results reveal (Table 15.14) that bandage PB5 has the lowest tear strength followed by PB1. It will be observed that there is no published benchmark or standard for tear strength that an ideal padding bandage should meet. Experience has shown that the bandage which possesses tear strength of 5000 mN or less, tested by using Elmendorf tear instrument, can be torn easily, by hand.

Relationship between demand absorption and wicking

It is a common practice in hospitals that venous leg ulcers are treated by a combination of wound dressing, under cast padding and compression bandage which are either a two-layer or multi-layer system depending on the severity of ulcers. The highly absorbent wound dressing should be used to absorb exudate and other body fluids from the ulcer. The padding bandage that is wrapped on the wound dressing should also to be highly absorbent to accommodate leakage of exudate from the wound.

It is observed that the absorptions of PB1, PB2, PB5, PB6, PB8 and PB10 are high and almost similar. On the other hand, PB3, PB5 and PB7 wicked high amounts of fluid. Similarly the rate of absorption of all the bandages is satisfactory except PB3, PB6, PBS and PB10. The bandages PB4, PB7 and PB9 possessed higher rates of wicking.

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Pressure distribution of commercial bandages

Padding bandage is applied beneath the compression bandage. The degree of pressure that is induced into the leg by the compression bandage is of major importance. It has been demonstrated that too high a pressure on the leg not only leads to further complications of the venous system but also promotes arterial disease. In contrast, inadequate pressure cannot help to heal the venous ulcers. Even if the compression bandage is applied at the correct tension it is probable that excessive pressure will be generated over the bony prominences of the leg. Therefore there is a need to distribute the pressure equally and uniformly at all points of the lower limb and this can be achieved by applying an effective padding layer around the leg below the compression bandage.

It is observed that none of the bandages provide uniform pressure distribution up to 60 mmHg. However PB5, PB6 and PB8 did distribute the applied pressure evenly only up to around 7 mmHg and the efficiency of the even pressure distribution degrades thereafter. It is vital that an ideal padding bandage should dissipate the pressure between 30 and 40 mmHg, exerted by high compression bandage, uniformly around the limb.

Adhesive bandages

An adhesive bandage usually has an absorbent pad (often medicated with antiseptic) covered by woven fabric, plastic, or latex rubber which has an adhesive. The plaster is applied such that the pad covers the wound (but does not stick to the wound), and the fabric or plastic sticks to the surrounding skin to hold the dressing in place and prevent dirt from entering the wound. Some newer plasters also contain woven strands of silver fibre, used to speed healing and minimize scarring. Adhesive bandages are generally applied to provide support rather than compression. The adhesive coating helps to ensure that the bandages do not slip or become displaced. These bandages are commonly used for strapping purposes and to provide support or compression in the treatment of sprains and strains. Diachylon adhesive bandage, more commonly known as Lestreflex, is often used in place of bandages coated with a zinc oxide adhesive where skin sensitivity reactions are present or suspected [163].

Adhesive bandages are characterized by their construction of two basic components, the adhesive coated backing material and the wound covering pad material. While such bandages are effective and desirable products, the assembly of the component materials during production results in increased manufacturing and inventory costs. In addition, the packaging of individual bandages requires additional handling and materials which further increases manufacturing costs. The bandage material is preferably
a heat-bondable, absorbent, nonwoven fabric which provides loft and absorbency in the pad area, while the single thickness wing portions of the bandage and the wrapper portion are heat calendered to provide a dense, sheet-like material. The bandage material is preferably a low density, highly absorbent, thermal bonded nonwoven fabric comprising absorbent fibres and staple length polyester-polyethylene conjugate fibres. These nonwoven fabrics are produced by a process which includes producing a web comprising absorbent fibres and staple length polyester/polyethylene conjugate fibres; subjecting the web to a temperature sufficient to fuse the lower melting component of the conjugate fibres without fusing the higher melting component while maintaining the web under little or no compression; and cooling the web to resolidify the lower melting component of the conjugate fibres, thereby forming a nonwoven fabric bonded at sites where the conjugate fibres touch each other and adjacent absorbent fibres.

A particularly preferred nonwoven fabric is a laminate comprising a core of a mixture of short-length natural cellulose fibres and staple length polyester/polyethylene conjugate fibres, and a lightweight veneer of heat-fusible fibres on each surface of the core. The composite web is passed through a through-air heater to fuse the lower melting component of the conjugate fibres while maintaining the fibrous integrity of these fibres, and to fuse or soften the surfaces of the heat-fusible fibres in the two outer veneers. As the material emerges from the heater and cools, the fused surfaces of the lower melting component of the conjugate fibres, i.e. the polyethylene, solidify, and bonds form where these surfaces touch each other and other fibres. Absorbent fibres employed in such thermal-bonded, nonwoven fabrics include rayon staple fibres, cotton fibres, short length natural cellulose fibres such as wood pulp fibres and cotton linters, and mixtures thereof [163].

Light support bandages

Light support bandages are also called short or minimal stretch bandages. They include the familiar crepe-type products of the British Pharmacopoeia together with numerous ‘non-official’ variations of these bandages, which are manufactured from cotton or cotton and viscose, and which show considerable variability in performance. They are used to prevent the formation of oedema and give support in the management of mild sprains and strains. Compared with the compression bandages, light support or minimal stretch bandages have limited extensibility and elasticity, and tend to ‘lock out’ at relatively low levels of extension. This feature enables them to be applied firmly over a joint to give support without generating significant levels of pressure. Short stretch bandages have also been used for the
treatment of venous leg ulcers. When applied at full extension, they form an inelastic covering to the leg, which tends to resist any change in the geometry of the calf muscle during exercise, thereby increasing surface pressure in a cyclical fashion and enhancing the action of the calf muscle pump [164].

Similarly, when an individual moves from a supine to a standing position or sits with the legs dependent, blood collects in the vessels and sinuses of the lower leg under the influence of gravity, causing the volume of the leg to increase. If this is associated with the formation of oedema, leg volume will increase still further. A short stretch bandage applied at full stretch, with the legs elevated, will tend to resist a change in volume as the legs are placed in a dependent position. This restriction will result in a significant rise in sub-bandage pressure, the degree of which will be determined by whether the subject is sitting or standing. Because short stretch bandages have limited elasticity, they are likely to be less effective than high compression bandages at reducing existing oedema as they lack the ability to ‘follow in’ as a limb reduces in circumference. They may, however, offer some advantages in the treatment of venous ulcers where a degree of arterial impairment is known or suspected as the low residual pressures will be less likely to compromise arterial inflow. Examples of light support bandages (including short stretch bandages) include Elastocrepe, Leukocrepe, Lenkelast and Comprilan.

Retention bandages

Retention bandages are used to retain dressings or other wound contact materials in position. They should not be used to apply pressure and are therefore totally unsuited for use in the management of leg ulcers or for the control of oedema. A number of different types of retention bandages are available. The first ‘retention bandage’ was White Open Wove (WOW), a rigid fabric with extremely limited conformability that is available in a range of widths. Because of the poor performance characteristics of WOW two softer more retentive bandages were introduced. These are Kling and Crinx, the cotton bandages of the British Pharmacopoiea. These lightweight cotton products have very little elasticity but are sufficiently extensible to give them a useful degree of conformability.

The ideal retention bandage should have a long shallow extensibility curve so that small changes in limb circumference will not significantly increase sub-bandage pressure, which in any event should not normally exceed a few millimetres of mercury. More recently lightweight woven and knitted bandages have been introduced which contain elastomeric yarns. These bandages, sometimes called ‘contour’ or ‘conforming stretch’ bandages, are often cheaper than the original cotton products and are said to
be easier to use. Many different types are available which include Slinky, Stayform, Tensofix, J Form and J Fast [164].

A new film dressing called Omiderm consists of a thin, flexible, transparent membrane of 40 microns thick. It is manufactured from polyurethane which has been chemically modified by the addition of hydrophilic monomers such as acrylamide and hydroxyethyl methacrylate. In the dry state the film is relatively inelastic but when brought into contact with wound exudates or aqueous solutions, it absorbs water and changes its physical properties, becoming highly conformable and elastic. Unlike traditional semi-permeable film dressings such as Opsite, Tegaderm and Bioclusive, Omiderm is not coated with adhesive but nevertheless will adhere to a moist wound without the need for additional fixative agents or sutures. The film is easily removed without causing pain or trauma but if left undisturbed it will separate spontaneously from a healed wound once epithelial cover is achieved. The film itself is highly permeable to moisture vapour, at least 20 times more permeable than the traditional film dressings, and is thus able to cope with the exudates produced from all but the most heavily exuding wounds. Omiderm is also significantly more permeable to oxygen than standard polyurethane film dressings. However the permeability of the film is not limited to water and oxygen. Antimicrobial agents such as silver sulphadiazine, povidone iodine and chlorhexidine gluconate can pass through the film onto the wound beneath if applied to the outer surface of the dressing in the form of a cream or as an aqueous solution absorbed on several layers of gauze. It has been suggested that this technique may be used to administer topical antimicrobial agents to wounds dressed with Omiderm to prevent infection or lower the bacterial count. Omiderm is recommended for the treatment of donor sites, dermabrasions and partial thickness burns and may also be used as a temporary dressing on full thickness wounds at the discretion of the medical officer in charge. It is not currently recommended for use on infected or dirty wounds [165].

15.3 Textiles for hygiene products

An increasing improvement in qualitative standards of human lifestyles has brought a greater sense of comfort and cleanliness. People are more and more looking for fresh public living surroundings and a higher level of hygiene in home areas. A wide class of micro-organisms coexists in a natural equilibrium with human body and living environments, but a rapid and uncontrolled multiplication of microbes can seriously compromise the hygienic and healthy personal standards. Because of their capillary spread in human living spaces, textiles have been involved in this research on improving the quality of hygienic living conditions. Many efforts have been
performed by the textile industry with two goals: the protection of the living environments and the textile fibres from an uncontrolled proliferation of microorganisms like bacteria [166]. Fabric treatments imparting bactericidal characteristics are highly desired by apparel, home, furnishing, and medical textiles. However, conventional processes used to impart such characteristics have a major drawback. That is, these effects are not permanent and the properties of the material may be altered. This problem has resulted in research efforts to develop durable treatments.

15.3.1 Antimicrobial textiles

Antimicrobial agents can be applied to the textile substrates by exhaust, pad-dry-cure, coating, spray and foam techniques. The substances can also be applied by directly adding into the fibre spinning dope. It is claimed that the commercial agents can be applied online during dyeing and finishing operations. The fibres derived from synthetic with built-in antimicrobial properties are listed in Table 15.15. We will discuss the following processes: blending, coating, finishing, chemical modification and grafting to improve the antimicrobial property of the polyester.

**Blending**

Incorporation of bioactive agents including antimicrobial agents into polymers by blending has been commercially applied in surgical implants and other biomedical devices. This is a pretreatment technology, where the antimicrobial agent is introduced during the processing stages. For this, the additive characteristics have to be compatible with blending conditions, e.g. particle diameter, heat stability, chemical stability and effect on fibre quality [166].

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Company</th>
<th>Brand</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyester</td>
<td>Trevira, Montefibre, Brilen</td>
<td>Trevira Bioactive, Terital</td>
</tr>
<tr>
<td>Polyacrylic</td>
<td>Accordis, Sterling</td>
<td>SANIWEAR, Bacterbril</td>
</tr>
<tr>
<td>Polyamide</td>
<td>Kaneba, R-STAT, Nylstar</td>
<td>Amicor, Biofresh, Livefresh, R-STAT, Meryl</td>
</tr>
<tr>
<td>Polypropylene</td>
<td>Asota</td>
<td>Asota AM Sanitary</td>
</tr>
<tr>
<td>Polyvinyl chloride</td>
<td>Rhovyl</td>
<td>Rhovyl's as Antimicrobial</td>
</tr>
<tr>
<td>Regenerated cellulose</td>
<td>Zimmer AG</td>
<td>Sea Cell Activated</td>
</tr>
</tbody>
</table>

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Silver–zirconium based antimicrobial agents have also been widely used for imparting long lasting antimicrobial properties [168, 169]. Wash fast antibacterial synthetic fibres like polyester were prepared by mixing metal ion supported inorganic compounds with fibre forming polymers. Polyethylene terephthalate and 2% Ag ion supported zirconium phosphate were melt spun to give antibacterial fibres exhibiting *Pneumonia bacilli* extinction to about 99% and 88% after 10 and 250 washings, respectively [170]. In another method, silver ions containing Zeomic show antimicrobial effects against a wider spectrum of microorganisms including Gram-negative bacteria such as *Escherichia coli* and *Pseudomonas aeruginosa*; Gram-positive bacteria such as *Staphylococcus aureus* and MRSA; and fungi such as *Aspergillus niger* and *Penicillium nigricans* (Table 15.16) [171].

Different bactericides like Cu, Ag, pyridine chloride, ethonium rivanol and kinamycin monosulphate have been used in polyester fibres where Ag and kinamycin monosulphate were observed to be active against gram-positive and gram-negative bacteria [172].

### Table 15.16 Antimicrobial effect of polyester cloth containing Zeomic [171]

<table>
<thead>
<tr>
<th>Tested bacteria</th>
<th>Test samples</th>
<th>Number of bacteria per cloth after contact</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0hr</td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>1</td>
<td>$3.6 \times 10^5$</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>$3.6 \times 10^5$</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>$3.6 \times 10^5$</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>1</td>
<td>$1.3 \times 10^5$</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>$1.3 \times 10^5$</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>$1.3 \times 10^5$</td>
</tr>
<tr>
<td>MRSA</td>
<td>1</td>
<td>$9.8 \times 10^4$</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>$9.8 \times 10^4$</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>$9.8 \times 10^4$</td>
</tr>
<tr>
<td>MRSE</td>
<td>1</td>
<td>$1.8 \times 10^4$</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>$1.8 \times 10^4$</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>$1.8 \times 10^4$</td>
</tr>
<tr>
<td><em>Trichophyton mentagrophytes</em></td>
<td>1</td>
<td>$3.5 \times 10^4$</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>$3.5 \times 10^4$</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>$3.5 \times 10^4$</td>
</tr>
</tbody>
</table>

Coating

In post-processing technology, the most common approach for applying the antibacterial agent is the coating. Antimicrobial agents that cannot
tolerate the temperature during polymer processing are often coated onto
the material after fabrication. The antimicrobial agents are linked to the
surface through physical bonds or anchored by cross-linking on the fibre.
Since most of these antimicrobial agents are water soluble, they are weakly
anchored onto the fibre surface so that it has to be constantly reapplied
[172].

An approach for antimicrobial coating of polyester is by using silk
sericin. Silk sericin is a natural macromolecular protein derived from silkworm, *Bombyx mori*. Sericin protein can be cross-linked, copolymerized,
and blended with other macromolecular materials, especially artificial
polymers, to produce materials with improved properties. The protein is
also used as an improving reagent or a coating material for natural and
artificial fibres, fabrics, and articles. Polyester fibres have micro-pores of
0.001–10 mm diameter. The sericin molecule can be introduced into these
micropores and cross-linked. Sericin-modified polyester fibre is obtained
by cross-linking with glyceryl polyglycidyl ether and diethylene triamine.
The sericin-modified polyester fibre can be more than five times as hygro-
scopic as untreated polyesters and more than 85% of initial hygroscopicity
remains after 50 washes [173].

*Finishing*

The following requirements need to be satisfied to obtain maximum ben-
efits out of the finish:

- Durability to washing, dry cleaning and hot pressing
- Selective activity to undesirable microorganisms
- Should not produce harmful effects to the manufacturer, user and the
  environment
- Compatibility with the chemical processes
- Easy method of application
- No deterioration of fabric quality
- Resistant to body fluids
- Resistant to disinfections/sterilization.

Hydrolyzed PET treated with 10% w/v seed extract (without any cross-
linking agent) showed antimicrobial activity of 89% and 32% even after
one washing against *Bacillus subtilis* and *Protues vulgaris* respectively.
This is may be due to more physical and hydrophobic interaction between
the hydrolyzed PET and the neem seed extract. The lesser antimicrobial
activity of neem seed extract against gram-negative bacteria is again proved
here.

From the Table 15.17, it is observed that antimicrobial activity of 84%
has been obtained after one washing using 10% neemazal technical (seed
Table 15.17 Antibacterial activity of neem seed extract (10%) treated PET fabric against *Bacillus subtilis* [174]

<table>
<thead>
<tr>
<th>Samples</th>
<th>Tested bacteria</th>
<th>Temperature (˚C)</th>
<th>Bacillus subtilis</th>
<th>Antibacterial activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Drying</td>
<td>Curing</td>
<td>Colony forming units (cfu)/ml</td>
</tr>
<tr>
<td>Original PET (Control-I)</td>
<td>–</td>
<td>–</td>
<td>335 × 10⁵</td>
<td>–</td>
</tr>
<tr>
<td>PET hydrolysed by 20% NaOH (Control-II)</td>
<td>–</td>
<td>–</td>
<td>340 × 10⁵</td>
<td>–</td>
</tr>
<tr>
<td>PET hydrolysed by 20% NaOH + C/L (citric acid Na₃PO₄H₂O₂), 1 washed (Control-III)</td>
<td>85</td>
<td>180</td>
<td>300 × 10⁵</td>
<td>10</td>
</tr>
<tr>
<td>PET hydrolysed by 20% NaOH + C/L (citric acid Na₃PO₄H₂O₂), 1 washed</td>
<td>85</td>
<td>180</td>
<td>55 × 10⁵</td>
<td>84</td>
</tr>
<tr>
<td>PET hydrolysed by 20% NaOH, unwashed</td>
<td>85</td>
<td>–</td>
<td>15 × 10⁵</td>
<td>95.5</td>
</tr>
<tr>
<td>PET hydrolysed by 20% NaOH, 1 washed</td>
<td>85</td>
<td>–</td>
<td>37 × 10⁵</td>
<td>89</td>
</tr>
</tbody>
</table>

extract) along with citric acid as a cross-linking agent. But one remarkable observation was that the fabric becomes yellowish to some extent when using citric acid as crosslinking agent [174]. *Table 15.18* shows the bacterial reduction against *S. aureus* after 30 minutes of cultivation. Researchers calculated the average bacterial reductions of each silver size. Average bacterial reductions of silver colloids explain why smaller sized colloidal silver has a better antibacterial efficacy than the others. Accordingly, 11.6 nm sized silver colloids had the best bacteriostasis, 99.9%. The growth of bacterial colonies was absolutely inhibited by only 10 ppm colloidal silver when the mean diameter of silver was 2–5 nm instead of 11.6 nm. Consequently, the smaller particle sizes had better antibacterial effects on silver-padded nonwoven fabrics. Higher concentrations of silver colloids have better bacteriostasis because bacterial reductions decreased when the silver concentrations in the pad bath decreased. These nonwoven fabrics are used for air filters or medical clothes [175].
A new class of N-halamine polymers has been synthesized. These polymers can be emulsified in water to produce coatings which, once chlorinated, act as contact disinfectants. The term ‘N-halamine’ herein signifies a molecule containing a nitrogen–halogen bond prepared by halogenations of an imide, amide or amine [176]. N-halamines were proven to be the suitable biocides that could provide desired antibacterial functions without causing much environmental concern [177]. The surfaces inactivate bacterial organisms efficiently, requiring relatively brief contact times of several minutes. The latexes can be formed by copolymerization of an N-halamine precursor monomer with other monomers in water with the aid of a surfactant, or by chemically grafting the N-halamine precursor monomer onto an emulsified polymer backbone, followed by chlorination. These coatings, once chlorinated, are effective at inactivating both Gram-negative and gram-positive bacterial organisms in relatively short contact times [176].

Antibiotics such as ciprofloxacin (Cipro), a quinolone antibiotic, can be directly incorporated onto the polymer using textile-dyeing technology, resulting in a sustained release of antibiotic over a period of time. Ethylenediamine (EDA) was used to bifunctionalize the Dacron material. Cipro covers a majority of the Gram-positive and negative bacteria that are encountered in a typical biomaterial infection, specifically Staphylococcus aureus.

A combination of specific surface characteristics can be incorporated into a single biomaterial. Functional groups are created with woven Dacron (Control) material via exposure to ethylenediamine (C-EDA). The antibiotic ciprofloxacin (Cipro) is then applied to the C-EDA material using

<table>
<thead>
<tr>
<th>Concentration ppm</th>
<th>Number of bacterial colonies /ml and % bacterial reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>11.6 nm</td>
</tr>
<tr>
<td>70</td>
<td>&lt;10</td>
</tr>
<tr>
<td></td>
<td>99.9%</td>
</tr>
<tr>
<td>100</td>
<td>&lt;10</td>
</tr>
<tr>
<td></td>
<td>99.9%</td>
</tr>
<tr>
<td>150</td>
<td>&lt;10</td>
</tr>
<tr>
<td></td>
<td>99.9%</td>
</tr>
<tr>
<td>Average</td>
<td>99.9%</td>
</tr>
</tbody>
</table>
pad/autoclave technique (C-EDA-AB) followed by surface immobilization of the coagulation cascade enzyme thrombin (C-EDA-AB-Thrombin). Antimicrobial activity by the C-EDA-AB surface has been observed to persist for five days compared with control and dipped controls. Thrombin surfaces had very high surface thrombin activity compared to nonspecifically bound thrombin and Cipro-dyed surfaces, respectively. Surface thrombus formation \textit{ex vivo} was evident after 1 min of exposure, with thrombus organization evident by 2.5 min. In contrast, C-EDA-AB and Control segments showed only blood protein adsorption on the fibres [178, 179].

\textit{Surface modification}

A variety of techniques are useful for chemical modification of polyester surfaces. These techniques include acid etching, X-ray irradiation, ultraviolet irradiation, electron beam bombardment, ozone treatment, and corona discharge and plasma treatments. A variety of polar groups are generated on the polymer surface as a result of these treatments. The generated peroxides and hydroxyl peroxides, in particular, are capable of initiating radical polymerization of vinyl monomers, resulting in surface grafted polymer chains [106].

\textit{Plasma grafting}

As in the case of plasma polymerization, plasma grafting began in the 1960s. In the 1960s and later in the 1970s, grafting experiments were mainly performed with acrylic acid and acrylic amide, preferably on fibres and textiles to enhance their wetability and receptivity for dyes. In the 1980s, especially with the work of Hirotsu, there was an increasing amount of interest in plasma grafting as it is desirable to modify polymers selectively for specific applications without losing much in terms of inherent characteristics [180]. The better the retention of the bulk properties, the more appropriate is the modification approach. In this respect, graft copolymers offer novel materials where the inherent polymer is represented by the backbone and the branches are formed by the grafted monomer with respective functionalities (Table 15.19) [181].

Chitosan possesses a good antibacterial property against various bacteria and fungi through ionic interaction at a cell surface, which eventually kills the cell [182]. PET texture was exposed to oxygen plasma glow discharge to produce peroxides on its surfaces. These peroxides were then used for the polymerization of acrylic acid (AA) in order to prepare a PET with carboxylic acid group (PET-A). Chitosan and quaternized chitosan (QC) were then coupled with the carboxyl groups on the PET-A to obtain
chitosan-grafted PET (PET-A-C) and QC-grafted PET (PET-A-QC), respectively (Figures 15.11 and 15.12).

The growth of *S. aureus* was not much influenced by contact with PET or AA-grafted PET. However, the growth of bacteria was significantly inhibited by contact with chitosan-grafted PET (62% in PET-A−-Cl+, 39% in PET-A-C, 59% in PET-A-QC). After 6 h of shaking the growth of bacteria was markedly inhibited by PET with ionically (86%) and covalently (75%) grafted chitosan and covalently grafted QC (83%). The PET-A−-Cl+ (86%) showed higher antibacterial activity than PET-A-C (75%). The high growth inhibition by PET-A-QC seems to be attributed to the quaternary ammonium ions of the grafted chitosan [183].

Nonwoven PET was activated by argon gas plasma. The plasma-activated substrate was immersed in the acrylamide (AAm) solution, which was sealed in a glass tube [184]. The antibacterial nonwoven PET can be

---

**Table 15.19** Functionalities imparted by monomers used in grafting [181]

<table>
<thead>
<tr>
<th>Monomer used in grafting</th>
<th>Functionality imparted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acrylic acid</td>
<td>Soil proofing, Antistatic properties,</td>
</tr>
<tr>
<td></td>
<td>Hydrophilicity</td>
</tr>
<tr>
<td>Maleic anhydride</td>
<td>Hydrophilicity</td>
</tr>
<tr>
<td></td>
<td>Adhesivity</td>
</tr>
<tr>
<td>Acrylamide</td>
<td>Antistatic properties</td>
</tr>
<tr>
<td>Polyethylene glycols (PEG)</td>
<td>Improving blood compatibility</td>
</tr>
</tbody>
</table>

15.11 Oxygen plasma treatment of PET and graft polymerization of acrylic acid (AA) on PET [183].
Polyesters and polyamides fabricated by the following methods. Method one is performed by immersing the AAm-grafted nonwoven substrate in AgNO₃ solution (concentration: 0.034–20.0 wt. %). Method two is to UV graft-polymerize the argon-plasma-activated substrate in the VQAS monomer aqueous solution (concentration: 10, 30, 50 wt. %). Method three is to immobilize the chitosan solution (solvent: 0.1 M acetic acid, chitosan concentration: 2 wt. %) with the glutaradehyde (GA) cross linking agent (CHO(CH₂)₃CHO, concentration: 1 wt. % GA/chitosan = 1:1) on the AAm-grafted substrate. The Ag⁺ ion treatment exhibits the best biocidal properties for the as-treated specimen shown in Figure 15.14. The preparation of the polyester–polyamide Ag-loaded textiles was carried out by RF-plasma and vacuum-UV (V-UV) surface activation followed by chemical reduction of silver salts. The rate of bacterial inactivation by the silver loaded textile was tested on Escherichia coli K-12 and showed long lasting residual effect (Figure 15.15).
15.14 Antibacterial results of PET nonwoven treated by different biocides: (a) original; (b) Ag$^+$ ions; (c) VQAS; (d) chitosan [184].

15.15 Inhibition of airborne bacterial growth by textile fabrics activated by RF-plasma as a function of the Ag loading [185].
Another approach for antimicrobial activity of nonwoven PET is to treat the plasma activation with argon and the subsequent UV-induced grafting polymerization of N-vinyl-2-pyrrolidone (NVP) to modify its surface hydrophilicity (NVP-g-PET nonwoven) [186]. The preliminary antibacterial assessment was determined qualitatively from the area from which *S. aureus* had been eradicated. The eradicated area was transparent. In comparison with the as-received nonwoven cloth, these three additive factor treatments improved the antibacterial properties of the PET nonwoven. The untreated PET nonwoven could not be anti-*S. aureus* growth; 28(b–d) could be anti-*S. aureus* growth because NVP integrating an iodine molecule acted as a biocidal agent. The improvement increased with an increase in the grafted amount of NVP on the surface of the PET nonwoven. It is apparent that NVP (additive AP and/or MBAAm) gave the best biocidal results [186].

\[ \gamma \text{-radiation} \]

Similar to plasma activation, \( \gamma \)-radiation also produces sufficient activation of the textile materials. Poly (ethylene terephthalate) fibers were treated with \(^{60}\text{Co-} \gamma \)-ray and grafted with acrylic acid (AA), followed by binding with chitosan (CS) via esterification. Afterwards, these CS-grafted fibres were immobilized with chondroitin-6-sulphate (ChS). The blood compatibility of PET was reduced by grafting with CS, while improved by immobilizing with ChS. The antibacterial activity of CS against *Staphylococcus aureus*, and *Pseudomonas aeruginosa* was retained after ChS-immobilization. After immobilizing ChS, the L929 fibroblasts cell proliferation was higher than CS-grafting PET fibres. The results indicate that by grafting with CS and immobilizing with ChS, PET fibres exhibit antibacterial activity and also improve the cell proliferation for fibroblast activity [187].

### 15.4 Intelligent textiles

The discovery of shape memory materials in the 1960s and intelligent polymeric gels in the 1970s was generally accepted as the birth of real smart materials. It was not before the late 1990s that intelligent materials were introduced in textiles. The first textile material that, in retroaction, was labelled as a smart textile was silk thread having a shape memory. Smart textiles can be described as textiles that are able to sense stimuli from the environment, to react to them and adapt to them by integration of functionalities in the textile structure. The stimulus as well as the response can have an electrical, thermal, chemical, magnetic or other origin [188].
In a recent report, thermosensitive material was prepared by radiation-induced graft copolymerization of monomers on PET fabrics. A binary mixture of N-isopropylacrylamide (NIPAAm) and acrylic acid (AA) was grafted on polyester fabric as a base material to introduce thermosensitive poly (N-isopropylacrylamide) chains having LCST close to 37°C in the membrane. The thermosensitive nature of the fabric was monitored by swelling at different temperature. The immobilization of tetracycline hydrochloride one model drug and its release characteristics at different temperatures were monitored.

The temperature sensitivity of equilibrium swelling in grafted samples was examined to determine their temperature dependent swelling and collapse process. No change in the swelling was observed for samples a, b and c since the mole fraction of NIPAAm grafted is very small as compared to AA. As a result, these samples don’t have a thermosensitive character. As the NIPAAm content in the copolymer increases, grafted samples start showing phase transition at specific temperatures. The first visible sign of the phase transition is noticed at 0.17 mole fraction of NIPAAm in the grafts. The LCST of the composition was 47.5°C. This temperature is known as lower critical solution temperature (LCST). The LCST of the graft copolymer decreases as mole fraction of NIPAAm increases (Figure 15.16). It is interesting to see that LCST of the fabric reaches 37.5°C for the NIPAAm fraction of 0.35 with grafted component. It may be mentioned that the grafting of pure NIPAAm exhibits LCST of 32.5°C as measured by DSC thermogram [189]. It may be stated that this composition of modified fabric may be useful for the drug delivery applications.

![Figure 15.16](image.png)

15.16 Variation of the equilibrium swelling with the temperature with degree of grafting (a) 9.6%, (b) 9.2%, (c) 8.6%, (d) 8.4%, (e) 8.4%, and (f) 7.7%.
One of the most attractive uses of PNIPAAm based copolymers is as drug carriers in which intelligent properties or auto-adjustables adjust to external temperature changes. It is important and practical to examine the T-HCl release data from the grafted samples having LCST (37.5°C) close to body temperature (37°C). The controlled release of T-HCl from PET fabric at different temperatures 20°C, 33°C and 37.5°C is shown in Figure 15.17. The drug loaded samples swelled at temperatures <LCST and deswelled at 37.5°C, in deionized water. Because the hygrogels shrink at 37.5°C, the T-HCl in the gels will be released due to the driving force of the volume change and concentration gradient of the drug. Hence, the amount released in the initial 10 minutes is highest at 37.5°C. Although at all the temperatures, samples exhibited very similar release profiles, their release rates and extents are different. The cumulative drug release during the 8 h study was 54% at 37.5°C, 14% at 33°C and 5% at 20°C. The observed initial burst release is possibly due to the drugs that were located near the graft copolymer surface. Since the concentration gradient is the driving force for the drug diffusion, a high drug concentration gradient between the copolymer surface and the release medium during the very early stage of contact leads to higher initial burst and fast release rate. Those drugs located near and at the surface could be released immediately from the graft copolymer to the surrounding medium as soon as the sample was placed into the distilled water [190].

A novel method of preparation of easily stripped off temporary wound dressing material is disclosed. In this process, the-N-isopropyl acrylamide (NIPAAm) monomer is successfully grafted on the non-woven polyester fabric by copolymerization. It is initiated by gamma-ray irradiation to activate the surface of the non-woven cloth. NIPAAm is then grafted onto
the surface of the non-woven cloth. The free radical or peroxide is produced by Co-60 gamma-ray, then grafted on the non-woven cloths. The lower critical solution temperature (LCST) in thermoresponsive poly-N-isopropylacrylamide (NIPAAm) is still retained after grafting. This will make the dressing cloth strip off easily without hurting the tissue. The material process is very simple and has medically applicable value [191].

15.5 Conclusion

Textile materials continue to serve an important function in the development of a wide range of medical and surgical products. Medical textiles are the products and constructions used as healthcare systems for example for first aid, clinical, hygiene implants and sutures for the huge medical market. The demands of the biomedical textile market have been growing worldwide at an annual rate of 4.6% because of their innovative features. Textile materials offer porosity and compliance which are often not exerted by other polymeric materials. Textile materials and products that have been engineered to meet particular needs are suitable for any medical and surgical application where a combination of strength, flexibility, and sometimes moisture and air permeability are required. Different forms of textile materials are used which include monofilament and multifilament yarns, woven, knitted, non-woven fabrics, and composite structures. The numbers of applications are huge and diverse, ranging from a single thread suture to the complex composite structures for bone replacement, and from the simple cleaning wipe to the advanced barrier fabrics used in operating theatres. Textile structures are particularly attractive to tissue engineering because of their ability to tailor a broad spectrum of scaffolds with a wide range of properties. There is no universal scaffold that meets the requirements of the various tissues of the human body. Further systematic study is necessary to design an optimal scaffold for each tissue application. Recent advances include the development of polylactic acid and polyglycolic acid fibres as structures for cell growth, temporary bioresorbable textile supports for growing human organic tissue such as bladder reconstruction, tissue engineering of vascular grafts, etc, and the development of smart fibres, based on naturally occurring polymers and also on non-animal-based protein fibres and structures for the treatment of wounds and ulcers.

Surgical implantation of these materials is encountered with both thrombosis and inflammation at the site of injury. These processes are related and both contribute to the healing of tissue into and around the material. Therefore, the main requirement of the textile material is bioreceptivity and biocompatibility at the application site in human beings. For this
requirement, it is necessary to modify the materials before using them in biomedical engineering. The various approaches to develop functional biotextiles are blending, coating, chemical treatments, and graft polymerization for making the surface bioreceptive and biocompatible. The introduction of new materials, the improvement in production techniques and fiber properties, and the use of more accurate and comprehensive testing have all had significant influence on advancing fibres and fabrics for medical applications. As more is understood about medical textiles, there is every reason to believe that a host of valuable and innovative products will emerge in the future with multidirectional growth in human healthcare systems.

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16.1 Introduction

This chapter looks at the technical, textile driven, design development of performance sportswear as a prime example of innovative applications for polyester and polyamide fibres. It considers the technical, aesthetic and cultural demands of the identified end-user as the starting point in informing the selection and positioning of specific textile assemblies, and trims, within an ergonomically designed clothing system. It looks at the sports ‘layering system’, traditionally tried and tested in military combat wear, as an inter-dependent combination of base-layer, mid-insulation layer and outer layer.

Until the emergence of man-made fibres, sports practitioners were often weighed down by military surplus clothing as well as by their heavy boots and accessories. The cut of early woven man-made fibre outer garments was necessarily voluminous, to provide ventilation prior to the introduction of breathable coatings, and to accommodate extreme movement. The silhouette of functional sportswear began to change in the late 1960s and 1970s, with the emergence of elastomeric yarns and waterproof, breathable fabric assemblies. Styling has become increasingly streamlined and sympathetic to the ergonomics of movement and the predominant posture for a given sport.

The needs of the body, in relation to the demands of the sport, provide a focus for the selection of an appropriate mix of fibre and yarn properties, woven, knitted and non-woven fabric constructions, assemblies and finishes. The characteristics of the textiles directly influence the cut, fit, and handle related to the specification of garment detail and construction methods to address protection, movement and posture for a breadth of different activities. The participation level and the varying degrees of expertise of the wearers, as well as the duration of the activity, or sequence of events, will influence the textile specification. Environmental and seasonal aspects of a sport, often taking place in extreme conditions, impact directly on the textile selection.
Modern sports clothing offers maximum performance, balanced with minimum weight, without sacrificing durability or comfort. Nylon and polyester fibres contribute to protective outerwear, insulating mid-layers, base layer and underwear as well as footwear and accessories. Sophisticated sportswear ranges demand design co-ordination throughout the clothing system with regard to colour, styling, proportion and trim. In the performance sportswear market textile fibres and innovative fabrics are invariably branded, or co-branded with apparel products, and often endorsed by internationally recognised, sponsored, sports ‘heroes’, such as Tiger Woods promoting Nike products.¹

16.1.1 The emergence of the sports ‘layering system’

The sports ‘layering system’ has evolved from military combat dress to address the selection of appropriate combinations of technical textiles in clothing to protect the body for a range of activities in contrasting environmental and climatic conditions. The clothing system is normally made up of an inner ‘base layer’, insulating ‘mid-layer’ and a protective ‘outer layer’ or ‘hard shell’. A new hybrid category has come into usage, that combines outer and mid-layer functionality promoted as the ‘soft shell’. In addition intimate apparel demands technical textile assemblies to offer support and to move with the body without chaffing. Personal protective garments or inserts, for contact sports or for extreme sports in hazardous environments, are also incorporated into the layering system.

Intimate apparel

Intimate apparel has been an early adopter of technical textiles and novel garment manufacturing processes since the introduction of Spandex for corsets in the 1960s. To achieve comfort, good fit and support, for both sport and fashion, lingerie and corsetry uses a breadth of woven, warp and weft knit, lace and non-woven materials in a range of fibre and yarn combinations. Weft knits offer more stretch around the body while warp knits are more stable in providing support. Warp knits are also used for swimwear. A variety of garment construction methods rely on a high percentage of synthetic fibres, predominantly nylon and polyester, in combination with elastomers, for the moulding of components and for their suitability in relation to the replacement of traditional garment sewing with new joining and finishing techniques. These processes include bonding, ultrasonic and laser welding, digital embroidery, engineered knits and seam free knit technology. Integrated textile sensors are now emerging in sports bras for monitoring health and performance, for example the Numetrex product range.

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Base layer

The term ‘second skin’ is often used for base layer garments worn closest to the body. These garments are normally made of knitted textiles and made both by traditional ‘cut and sew’ garment construction methods and by circular knit construction. Lightweight warp and weft knitted fabrics are usually selected for ease of movement, providing stretch in their stitch construction to fit like a second skin. Base layer fabrics for sport are made from fibres and yarn combinations that wick perspiration away from the skin to prevent chilling, especially when stationary and in extremely cold conditions. Synthetic fibres promote moisture management and dry much quicker than cotton. Polyester has good wicking and fast drying properties, while nylon is more abrasion resistant. ‘Added value’ may be provided through a range of fibres and finishes that offer anti-microbial and anti-ultraviolet (UV) protection.

Mid-layer

The mid-‘insulation layer’ may be varied in thickness or bulk, to alter its ability to trap ‘still air’, or ‘dead air’, depending on the demands of the range of likely conditions. This layer should have compatible cut and styling, with that of the base and outer layer garments, to avoid impeding movement. The insulation layer may be made up of more than one garment; with examples such as jackets, smocks, gilets, traditional knitwear, felted garments and quilted woven textile assemblies incorporating down and synthetic waddings. The most important innovation in insulating textiles for sport, however, has been the development of knit structure fleece fabrics. These fabrics are primarily weft knit and polyester based although nylon adaptations and warp knit structures are also in use. While sophisticated fleece constructions are finely tuned to particular specifications for performance sport, mass-market derivations embrace men’s, women’s and children’s wear ranges in the high street for both sport and everyday life. A recent innovation has been based on the concept of the trapping of still air in a garment that can be filled with air, via a network of internal chambers, and the level of insulation regulated via a tube attached to the inside of the vest to keep the wearer warm.\(^2\)

Outer layer

The usual requirement for an outer shell garment is to be super lightweight with minimum bulk for easy storage.\(^3\) The ‘shell’, or protective layer, is selected to provide the most appropriate balance of windproof and ‘waterproofness’ versus ‘breathability’ and protection for the specified range of
activities. The wind chill factor can lower the ambient temperature dramatically and threaten the clothing microclimate. The outer shell, traditionally also known as ‘hard shell’, is intended to protect and maintain the function the whole ‘layering system’. ‘Soft shells’ provide a relatively new generation of outer layer garments that have emerged as ‘all-rounders’ for winter and summer. These are softer, bulkier and warmer than hard shell garments and often with stretch for comfort.

Personal protection

For certain contact sports, or for those carried out in hostile environments, the governing bodies’ rules have standard requirements for health and safety that directly impinge on design. For extreme sports, such as motor biking and, more recently, snowboarding, textile fibres are being incorporated into a sophisticated mix of spacer fabrics and heavy meshes in exciting ergonomic designs.

Other market opportunities that have the potential to benefit from lightweight protection and the comfort and easy care attributes of polyester and polyamide fibrous materials include corporate wear, travel wear and inclusive design, with particular relevance in the promotion of health and wellness. A well-designed, versatile, ‘layering system’, has the potential to address the functional demands of the modern, global, 24-hour society, mixing work, relaxation and everyday activities for an inclusive audience.

16.2 Fibre developments and characteristics

The marketing hype, associated with the branding of fibres, escalated towards the end of the twentieth century with Dupont branding spandex as Lycra in 1960s, W. L. Gore introducing waterproof breathable laminates as Gore-Tex in the late 1970s and ICI re-launching nylon as Tactel in 1983. Performance sportswear is now brought to market through the promotion of internationally recognized brands with clothing products often co-promoted through partnerships with fibre and textile producers. Fibres are promoted through a marketing ‘story’, in a plethora of point of sale (POS) material, often linked to the performance of top athletes. In this new century there is a growing concern for environmental issues associated with the textile chain, as well as an emphasis on textile attributes that can promote health and wellness. Although it is almost impossible to ignore the claims of the brands, with their changing ownership, it is important for designers, and those in the product development team, to have an appropriate level of understanding of generic fibres, their production, properties and characteristics.
16.1 The application of technical fibres and fabrics in the sports layering system.

16.2.1 Fibre and yarn extrusion

In man-made fibre production the fibre is extruded in the form of continuous multifilament flat or straight filaments referred to as continuous multifilament flat yarns. These thermoplastic filaments may be permanently distorted, under the influence of heat, to produce bulky or textured
filament yarns. These yarns may be bulky and elastic or simply bulky. Fibre producers introduce many interesting variations of nylon and polyester fibres in the extrusion process, by altering the shape of the spinneret holes, to produce multi-channel filaments in a range of multilobal shapes. The handle and light reflecting properties of a fibre may be modified by changing the cross-sectional shape, for example trilobal, pentalobal and octalobal filaments. Hollow channels may be introduced into synthetic fibres to increase thermal insulation. Ultra-fine microfibres give enhanced flexibility, wicking and water repellency coupled with water vapour permeability. Moisture wicking may be enhanced by the use of appropriate hydrophilic surface finishes, by the incorporation of ‘water attracting’ co-polymers. Other finishing techniques such as soil release, flame retardant and antimicrobial attributes are available.

Some examples of well-promoted polyamide and polyester branded fibres, adopted within the sports layering system, are as follows. The nylon fibre Tactel®, originally launched by ICI in 1983, has since passed from DuPont to Invista. A current product, licensed to Nilit, is Tactel® multisoft with a round or trilobal cross-section. AdvanSA’s moisture management polyester fibre, Coolmax®, has both four and six channel variations, with an increased surface area to promote moisture wicking. AdvanSA’s polyester hollow fibre is marketed as Thermolite® to provide added insulation. Co-polymers may be introduced to give dye variant fibres, with two different colours in one dying process or tone-on-tone effects such as Nilit’s proprietary brand Sensil® Duelle or their licensed brand, Tactel® Strata. Fibre properties may be improved through further modifications to the basic man-made fibre extrusion processes. High-tenacity nylon fibre, such as Tactel HT, that is drawn after extrusion, is known to be lighter and more abrasion resistant than other nylon fabrics. The nylon fibre, Cordura® is said to be twice as durable as standard nylon and three times as strong as polyester.

16.2.2 Micro-encapsulation

Micro-encapsulation or impregnation techniques may be used to introduce anti-microbial properties that reduce odour formation and the risk of infection. Antibacterial agents can be added to synthetic fibres in the spinning process. ‘Sanitized’ silver yarn has permanent bacteriostatic properties due to the incorporation of sanitized silver nano-particles in the PA6 polymer mix before spinning which prevents the proliferation of bacteria and the build up of unpleasant odours in garments – even if washed frequently. Because of its PA6 polymer matrix it will not yellow and takes up dyes exceptionally well.® Another example is the newly launched ‘Cocona’ fibre, the result of a partnership between Burlington Worldwide and Trap Tek, that has activated carbon particles, from heated coconut husks, per-
manently embedded into polyester fibre. This provides a porous structure that traps odours, promotes the evaporative cooling of perspiration, and offers protection from harmful solar, ultraviolet radiation.

Another example of micro-encapsulation is in Phase Change Materials (PCMs). ‘Outlast’, USA, is a branded product, originally developed for the garments of NASA astronauts, that buffers temperature swings, absorbing excess heat when the wearer is active and releasing heat when the wearer begins to cool down. This is described as a ‘microencapsulated latent heat storer, based on paraffin wax’. This is ‘the phenomenon that occurs when the wax changes from the solid to the liquid state. During this phase transition a large amount of thermal energy (latent heat) is consumed without the temperature of the material itself changing. The innovative, design led knitwear company, Falke, has incorporated Outlast ‘Adaptive Comfort’ phase change technology into polyamide fabrics for their Autumn/winter 2007/2008 ‘Ergonomic Sports Underwear Collection’ with styles marketed as ‘Athletic Warm’ and ‘Comfort Warm’ that aim to provide ‘a balanced body climate’.

16.2.3 Fibre blends

All fibres have good, fair and poor characteristics relating to a particular end-use. Pharr Yarns, maintains that ‘the market is driven by the properties the end-user wants’. This US spinner steers their clients towards an appropriate blend as ‘a yarn made of 100% of any fibre will not accomplish all that a customer wants.’ Fibre blending allows manufacturers to combine fibres so that good qualities are emphasised and poor qualities minimised. Blending may be carried out at any stage prior to the spinning operation. The earlier the fibres are blended in processing, the better the blend. Blending may be carried out to give colour effects, to obtain better texture, to alter handle or fabric appearance and to improve performance. Special categories of blended yarn have been developed to exploit particular fibre characteristics. Some yarns have two types of continuous filament intermingled within one continuous filament yarn. Others have two types of polymer within the same filament. A blend of nylon fibre and silver-coated nylon fibre, known as X-static, is an effective sustained-release antibacterial agent. Composite yarns have two or more elements one of which is a continuous filament. Core spun elastane has a central core of elastane often covered with polyester or nylon wrapping yarns.

16.2.4 Stretch

Ease of movement has become a highly desirable fabric property not only in performance sportswear, but also in everyday clothes. Stretch fabrics
enhance comfort and fit and ease the putting on / taking off of garments and provide good shape retention. High power elastomeric stretch can also provide a valuable supportive function, applying compression to injured limbs, muscular support or support to a bust line. There are three types of stretch yarns. ‘Elastomerics’ are those where elastic recovery is a fibre property. Although elastomerics may be used bare, they are often covered with a non-elastic yarn such as nylon or polyester. Crepe or ‘over spun’ yarns are those where the high degree of twist in the spinning process makes the yarn pull back on itself. Texturized yarns are those whereby a false twist is manufactured during the production of synthetic yarns.

Other materials have highly elastic properties, e.g. special polyesters such as polybutylene terephthalate (PBT). These materials are used in swimwear, support garments such as athletics garments, skiwear, leotards and increasingly in comfort stretch general clothing. PBT is less susceptible to degeneration in chlorine or sunlight. An elastic bicomponent polyester fibre, generic name ‘elasterell-p’, is a hybrid between elastane and textured polyester. It is not too elastic and therefore suitable for fabrics requiring moderate stretch or comfort stretch providing better-fit and good shape retention. With extraordinary tensile strength, it can withstand tough wear, extending garment life.

16.2.5 Sustainability

In the twenty-first century designers, and those involved in the textile related product development teams, should have an awareness of sustainable issues in relation to the selection/specification of textiles from fibre development to textile processing, finishing and after-care. This is particularly relevant for performance sportswear products where the users are normally concerned with health, wellness and care of the environment. As recycling becomes compulsory, so sustainability issues will increase in importance and the ‘end of life’ for technical garments must be considered. The outdoor clothing company Vaude, and partners, has pioneered ‘Eco-log’ whereby every component in the clothing system is made of polyester from waterproof weave with Sympatex, polyester based, membrane, to fleece mid-layer and wicking base layer. Polyester zips, studs, threads and trims complete the mix enabling the whole garment layering system to be recycled. The US outdoor company, Patagonia, has used recycled PET (polyethylene terephthalate) fleece fabrics for over a decade, initially from Wellman in the USA and now from Teijin in Japan.

Biodegradable, crop-based, polymers may be produced from corn starch, an abundant natural polymer, cellulose, as an alternative to fossil fuel synthetics. PLA (polylactose acid) fibre is said to ‘bridge the gap’ between natural polymers and synthetics, offering a unique combination of proper-
ties with the best attributes of natural and synthetic fibres. It is claimed that biodegradable polyester has the easy care properties of real polyester and the right technical credentials to replace polypropylene. Cargill Dow’s PLA fibre ‘NatureWorks’ is now extruded into a high-performance synthetic fibre, branded as Ingeo, claimed to be ‘the world’s first man-made fibre derived from 100% annually renewable resources’. Yarns are being developed, to include apparel applications, in both pure qualities and innovative blends. Strength and resilience are balanced with comfort, softness, drape and good moisture management characteristics. It may be used as fibre fill for products such as padded outerwear with superior loft that is said to feel more like down than a synthetic imitation.

Marks & Spencer (M&S), the major retail chain, announced ‘Plan A’ in January 2007, to provide leadership on the issue of sustainability over the subsequent five years and beyond. With regard to textile products, M&S plans to promote the use of sustainable raw materials including PLA and polyester made from recycled PET bottles. ‘Plan A’ includes the planned opening of a model ‘green factory’ in collaboration with a supplier. Recycling points will be introduced for packaging and in-store points for the consumer to be able to recycle clothing. The aim is that within five years the consumer will not need to throw away any M&S clothing waste after they have finished with it. There are also plans to start researching alternatives to clothing disposal, including donation, composting and recycling.

16.3 Design considerations
16.3.1 Technical garment design

The adoption of technical textiles in lightweight nylon and polyester fibres has directly driven design innovation in sports clothing in terms of comfort, performance and aesthetics. The demands of the body, for movement, moisture management, thermal regulation and protection may all be addressed through the application of modern technical textile developments with nylon and polyester fibres as major constituents. The versatility of the performance sportswear clothing system enables top athletes and recreational sports enthusiasts to carry out individual or mixed activities, over short or extended durations, without the need for heavy or bulky clothing and accessories that impede movement. In combination with technical performance attributes, polyester and nylon fabrics may provide sophisticated aesthetic appeal.

A slim fitting silhouette is now the norm as a result of innovation in breathable fabrics and stretch properties. This, in combination with ergonomic cut, avoids billowing when moving at speed, with the front hem of a jacket invariably hollowed out and the back lowered, relevant to the
predominant posture desired. For outdoor performance clothing, garments incorporate details such as optional foldaway hoods into collars and pockets for maps, phones and security items. Other garment details, which use a range of nylon and polyester textile constructions, include abrasion resistant articulated elbows, hook and loop cuff adjustment (commonly marketed as Velcro®), zips for closures and ventilation as well as shock cords, fleece facings, garment insulation and protective spacer fabrics.

The potential combinations of nylon and polyester fibres, and blends, in fabric structures for mid- and base-layer garments is infinite in providing a breadth different attributes. Strategic ‘zone construction’, or the placement of fabrics, in relation to the needs of the body, increases performance with relevance to the end use. Seam free and whole garment knitting technology is integral to the design of garments that adopt these all-in-one production techniques. Mesh structure inserts increase air flow for increased moisture wicking in high sweat areas, mixed fabric weights can be selected to enhance mobility, give protection and can add durability in areas of high abrasion. Traditional garment construction is done in two dimensions on a flat surface while new functional garment construction methods follow the three dimensional contours of the body to provide reduced friction, increased movement and protection and ergonomic shaping. Functional garment construction benefits from features such as gussets that reduce friction in key areas like underarm; raglan sleeves keep the shoulders free from seams that might rub; and articulated construction enables freedom of movement for elbows and knees. Additional shape and protection may be provided in the moulding of garment components, commonplace in intimate apparel garments, and for lightweight body armour.

Leading edge stitch free garment construction techniques are emerging throughout the technical textile driven performance sportswear layering system. Garment bonding and laser welding are stitch free joining technologies used in high end markets to provide clean, streamlined design both for woven and knitted textile constructions without inhibiting stretch. Flat seams with clean finishing enhance comfort inside the neck and around sensitive areas. Laser cut edges and pockets with bonded waterproof zips reduce weight in both hard and soft shell garments. Sonic stitching is used primarily for disposable garments such as forensic suits, medical end use and clean room garments. Performance sportswear has been an early adopter of novel manufacturing techniques that are gradually filtering through to mainstream fashion.

16.3.2 Analysing garment characteristics

When designing a performance sportswear ‘product’ there are many factors to consider. These should be fully explored and identified by starting with
an analysis of the intended end-use. The design development process should begin with identifying the ‘customer’ or market sector in relation to the specified sport or range of activities. The range of issues uncovered may then be prioritised to inform the garment design brief and help guide the subsequent product development.
Questions to be addressed may include the following. Is the garment or product intended for individual use or team wear and what is the level of expertise of the wearer(s). Is the garment incorporated in a layering system and, if so, what are the other layers? What are the demands of the likely environmental conditions, or range of locations, that may be potentially hazardous, as well as weather and seasonal considerations? What is the duration of the activity or sequence of events? The predicted life span of the product must be considered and a plan for its disposal. Will the material selection influence the type of garment construction?

Is the product to be in contact with the skin and what are the demands of the workload? Is the activity a contact or non-contact sport in terms of body protection? The ergonomics of movement is a major consideration in performance sportswear design. Predominant posture and extreme movement varies for different activities but there are generic considerations for performance sportswear that are not commonplace in fashion. For a breadth of sporting activities it is important to have extended arm lift, articulated elbows and knees and a lack of restriction across the back and in the leg-wear crotch area. Movement may be enhanced through the garment cut but also in the selection of appropriate fabric constructions, making use of bias cut and through the incorporation of elastomeric fibres and yarns.

The garment characteristics required, from both technical and aesthetic perspectives, will inform the drafting of a clear and effective design brief to guide decision making with regard to the most appropriate textile fibres, fabric structures and finishing choices for the chosen ‘customer’ or market. Nylon and polyester fibres are prevalent in a range of products throughout the sourcing and selection of functional fabrics, trims and components, from linings and laminates to knits, weaves, cords and zips, that enhance the comfort, functionality and appearance of highly detailed performance sportswear outer garments.

16.4 Textile selection

Nylon and polyester fibres constitute a breadth of different structures and assemblies for performance sportswear applications that include wovens, non-wovens, weft and warp knits, nets, meshes and narrow fabrics. Woven fabrics are generally strong, relatively rigid and hardwearing and tend to be used for outer layer sports garments. Woven constructions are also used for back-pack fabrics and their webbings and trims. Lightweight weaves and nets may be used as linings. Non-woven textiles are found in protective laminates and insulating waddings. Knitted textiles provide highly flexible structures throughout the clothing ‘layering system’ from base-layer, or ‘second skin’, to mid-insulation layers and to both the linings and main
substrate of many protective outer layer garments. Textile structures, with their branded fibres, may be loosely categorized within the different layers of the generic sports clothing system. The base layer is primarily concerned with moisture management, the mid-layer with insulation, and the outer layer with protection.

16.4.1 Textiles for moisture management

It is important to move moisture away from the skin and to maintain a dry microclimate as damp base layer garments can cool rapidly, be uncomfortable, and may cause post-exercise chill. This is vital in high activity winter sports which take place in cold climates by comparison with hot dry desert conditions where cooling is desirable. ‘During intensive training, an athlete’s clothing has to transport around 1.5 to 2.5 litres of moisture per hour.’ The breathability, heat and moisture management of textiles play a crucial role in supporting the physical and mental performance of the wearer.

Double face ‘denier gradient’ fabrics use a plaited technique that joins two different constructions to manage moisture through push-pull capillary action. The body’s heat pushes sweat and moisture vapour through the larger gradient fabric, next to the skin, towards the outer surface (face), and the micro-yarns on the face pull the moisture to the surface, absorbing and spreading it quickly to evaporate and leave a dry surface next to the skin. A dry inner surface feels more comfortable and reduces chaffing. AdvanSA and Lenzing have launched a new range of fabrics that have polyester Coolmax® to be worn against the skin, and absorbent, cellulosic Tencel fibre, with a natural touch, on the outer face of the fabric.

A close body fit enables faster moisture absorption to speed the wicking process. When close fitting garments are required, lightweight weft knits give maximum stretch around the body. Weft knitted fabrics have more stretch in the course, or horizontal direction of the fabric, than in the wale, or vertical direction, which has minimal stretch. Weft knit fabrics are relatively bulky and open in construction being air and water permeable providing good insulating properties as well as breathability. Weft knits are used in intimate apparel, base layer garments, swim wear, cycling wear, team wear as well as in open mesh garments/linings and fleeces. Fabric elasticity may be enhanced by the use of elastomeric fibres. Garment fit, for a breadth of figure types, is a key consideration for the embedding of wearable technologies in base layer garments to monitor the wearer’s performance for sport as well as general health and wellness.

Circular weft knit innovation is being led by the machine makers, such as Santoni and Stoll. Santoni ‘seamfree’ technology can be used to
engineer seamless structures to create garment fit and the placing of controlled ‘zones’ of different knit constructions, with jacquard patterning, surface dimensionality and the selection of different colours and yarn properties. Areas can be incorporated with three dimensional and open mesh stretch support to create shaping with very little sewing required. Engineered customization can be realized by integrating other technologies, such as a body scanner and performance yarns.

At the major sports trade fair, ISPO, Spring 2007, the Swedish company, Craft, presented seamfree ‘Pro Warm’ base layers that use ‘very lightweight hollow polyamide fibres, polyester and elastane in a complex 3D knit with seamlessly integrated climate zones. While the hollow fibres and spaces in between trap air, thus providing the insulation, the climate zones are said to guarantee excellent temperature management, keeping the body warm where it is needed, yet preventing overheating. It is also said to be super soft, providing a high level of moisture management, good fit and freedom of movement.

16.4.2 Textiles for insulation

Weft knitted fabrics in different variations of weight and bulk are prevalent in ‘mid-layer’ garments. They tend to be more extensible and elastic than woven fabrics. Fibre and fabric mixes for insulation require bulk and include fleece, or brushed pile fabric, fibre pile or sliver knit, sometimes known as fake fur, and other three-dimensional knitted spacer constructions. Insulating fibres may be shaped and/or hollow and often made from polyester to prevent absorption of moisture. ‘The invention of fleece changed the entire apparel, sports and outdoor market. It is even considered to be one of the top one hundred inventions of the 20th century.’ It is now worn as much in everyday wear, and for corporate workwear, as it is for performance sport for all age groups.

Mountain Hardware’s ‘Euro Stretch’ is a dense bi-elastic two sided fleece fabric that can be worn either way, allowing a smooth outside and plush inside, used in their ‘Expedition reversible T’. Some warp knitted constructions also have one smooth face and one textured loop or brushed face, so that, worn one way, garments trap air and offer extra warmth in cold conditions and when reversed offer cooling comfort. This reversible technology was developed by Nikwax, UK’s Paramo range, in their ‘Parameta S’ shirts.

As with base layer garments, knitted mid-layer fleece developments can be engineered to embrace different ‘zones’ of insulation in relation to the ergonomic design of the product and the requirement for protection, thermal regulation and moisture management relevant to the demands of different areas of the body. Seamless garment construction reduces any
chaffing as traditional joining is eliminated. The US outdoor company, Patagonia, developed a variable knit fleece with Malden Mills, the originators of ‘Polartec’ fabrics.

In addition to insulation in mid-layer garments, other technologies have been discussed that address thermal regulation such as phase change materials, that retain or release heat depending on workload. Outlast® molecules can be incorporated through fibre micro-encapsulation but also incorporated into the garment system as a laminate. Wearable electronics can be embedded in mid layers for products, such as fleeces, with battery-driven heating providing warmth to the kidney area.

16.4.3 Textiles for protection

Outer layer garments are generally made of woven constructions with coatings or laminates incorporated into two or three layer assemblies. These are also treated with water repellent finishes on the outer face. Hard shell outer fabrics are normally of lightweight nylon or polyester. Nylon fibre is used for its strength and abrasion resistance for more extreme end-use while polyester is selected for a softer handle and is more suited to recreational sport. Garments may have enhanced protection with areas of abrasion resistant fabrics such as nylon Cordura. These placements are often on the elbows or knees, across the shoulders or to reinforce an area that has regular use such as the side of a garment if carrying a snowboard. Lightweight nylon weaves can be strengthened with rip-stop weave structures. Polyester weaves may have enhanced softness and drape through sanded or emerised finishing.

There are various descriptors for protective outwear garments categorised by their fabric assembly. Within the ‘Hard Shell’ category, ‘two-layer’ garments are made from fabric that has an exposed coating or laminate on the inside, normally protected by a loose mesh lining, and ‘three layer’ garments have a ‘sandwich construction’ with an outer woven fabric, a laminate and a backing to protect the laminate, such as a fine single jersey or mesh. The outer shell design must incorporate appropriate ventilation as few textile assemblies cope with the moisture produced from extreme workload.

The new class of outerwear garment, the Soft shell, normally incorporates stretch for added comfort. Soft shells are often of a ‘sandwich’ construction of stretch woven outer, a laminate, and a fleece or brushed mesh inner backing. Some ‘soft shells’ have additional anti-abrasion ‘hard shell’ laminates on outer areas that need greater protection and reinforcement. Outer fabrics are often of nylon fibre for strength and inner fabrics are often polyester for softness. Successful designs combine a mix of material compositions and constructions, placed in positions appropriate to the
demands of the body, to achieve maximum comfort and performance. Sports companies adopt their own terminology in developing garments using garment construction techniques. ‘Body mapping’ is the terminology used by W. L. Gore to address the placement of textiles with a range of attributes to address the needs of different zones of the body, that may require extra padding, moisture absorption or heat insulation. For personal protection a stable warp knit structure is used to produce three-dimensional spacer fabrics in a two-layer ‘sandwich’ construction. Monofilament connecting yarns maintain the space between the layers.

Waterproofness and breathability continue to be major requirements for sports outerwear garments. The branded waterproof breathable laminate, ‘Sympatex’, is a polyester based membrane. Promotional material from the company claims that their product ‘Reflexion 111’ is a ‘membrane with optimised heat management providing efficient insulation in cold temperatures while also offering breathability in a light weight form. The compact hydrophilic membrane is thinner, lighter and more flexible (300% bi-stretch) than other membrane systems as well as being waterproof, windproof and recyclable. The non-porous surface is plasma treated and aluminised in a 4000 degree vacuum environment using an extremely thin 50-nanometre layer of aluminium. The laminate can be sewn, seam sealed or bonded. Fine continuous filament flat yarn warp knitted fabrics are normally used to protect the inner layer of laminates. Woven linings are used for more tailored elements and warp knit mesh fabrics are widely used for loose linings and pocket bags.

### 16.5 Future trends

Innovation in performance sportswear design continues to be driven by technical textiles, in particular the attributes of lightness in weight, stretch, moisture management, thermal regulation and protection. Increasing sophistication in garment cut and fit, enhancing movement and protection, is directly related to advances in synthetic fibre developments within novel textile structures, in tandem with new garment construction. We are now at the beginning of a new revolution in the textile and clothing industries as the disparate areas of technical textiles and performance clothing merge with the electronics industry and related sectors such as communications and health.

The embedding of smart attributes and electronics in textiles further enhances the comfort of clothing through the protection and maintenance of the clothing microclimate. Functionalities that include phase change and shape memory polymers, sensors that measure human vital signs, positioning and performance, heated textiles and self-cleaning nano finishes are being brought to market in sectors such as corporate workwear, health and
wellness products and in performance sportswear. Many of these textile innovations, for performance sports end-use, are embedded in textile assemblies that are largely constituted from polyester and polyamide fibres.

16.5.1 Biomimicry

Many examples of innovations in modern fibres and textile structures have resulted from the principles of biomimicry. Textile applications that improve functionality in sportswear, have drawn inspiration from lotus leaves, mimicked in self cleaning fabrics, and gecko feet mimicked in closures. Fir cones were one of the earliest examples cited as ‘a flexible response to moisture and adaptation to temperature change. When the weather is cold or wet they close up and when dry they open again.’ In Schoeller’s ‘C-change’ technology, a hydrophilic membrane is set to a particular temperature range and, as soon as higher temperatures or body warmth result in greater moisture being produced, the flexible polymer structure adjusts to allow water vapour to escape quickly to the outside air. When the body produces less heat energy and therefore less moisture, the polymer structure returns to its original condition.

16.5.2 Shape memory polymers

‘Shape memory polymers (SMPs) are smart materials that, as a result of external stimulus such as temperature, can change from a temporary deformed shape back to an original shape.’\textsuperscript{14} Shape memory polymers offer flexible lightweight impact protection for extreme sports. Dow Corning has an ‘Active Protective System’ described as ‘3-D spacer textile treated with special silicone coating which remains soft and flexible under normal use but hardens instantly upon impact’. This foam-based material immediately reverts back to being pliable material once the pressure is lifted.

An alternative protective technology is described as ‘Rate Sensitive Material’. The product ‘d3o’, is made up of free flowing molecules that provide a soft pliable material which reacts the moment impact occurs. Once the impact is over, the material immediately becomes flexible again. The molecules lock together to provide a shock absorbing system that may provide removable protectors in areas such as shoulders and elbows and knees. This product is also used in the relatively thin and stylish Reusch ‘Unshok glove’. This material both absorbs and distributes impact force with the reaction starting sooner and lasting longer than in rigid systems.
16.5.3 Wearable electronics

Conductive fibres are of particular interest in relation to smart textile applications and wearable technology, both in terms of the elimination of static, and also as conductive pathways for the connection of electronic components. Conductive materials include metal fibres as well as inherently conducting polymers (ICPs) and carbon fibres already being used in many applications for occupational work wear such as antistatic working, EMI (Electromagnetic Interference) shielding, heating and the transport of electrical signals. Some sportswear companies have already had great marketing success with wearable electronics from the simplest applications to complex systems. Smart textiles, with embedded technology, are emerging in a rapidly growing range of applications that include heat conductive textiles, textile switches and vital signs and performance monitoring devices.

Several UK-based companies have integrated electronics into sports clothing and textile based accessories. Companies, including Eleksen, EXO, Fibretronic and Peratech, have designed products that interface with software from Microsoft and iPod music players from Apple. Products are being developed with internationally recognised sports brands such as Berghaus, Burton, Nike, O’Neill, Quiksilver, Reusch, Rohan, Schoeffel, Spyder and The North Face. One of the most prominent technology providers in the sportswear market has been Eleksen who specialised in touch-sensitive interactive textiles for electronics interfaces. The core technology is the use of an electro-conductive fabric touch pad, ElekTex, for the creation of flexible, durable and rugged fabric touch screen interfaces. This technology has been employed in products that include interactive sports apparel, bags, back packs and communications controls in textile computer keyboards that can be rolled up into a compact space.

The UK outdoor clothing brand, Berghaus has developed the ‘Heatcell Gilet’, a fleece containing the EXO with a ‘Thermomesh’ heating system, edged with conductive tinned copper braiding, encapsulated in a nylon envelope, that spreads warmth around the torso for up to six hours. This type of climate control system is also used in the ‘StormRider’ heated vest for professional motorcyclists. Many of the well-known sports and outdoor brands are now employing technology and electronics specialists. Andreas Roepert, of Interactive Wear AG, advises that ‘Attention to the differences in the design and production processes, a joint project plan and timely cooperation between wearables and textile partners helps to fashion the design successfully and to optimise costs’ and that ‘well thought out production directives save some surprises’.
16.6 Sources of further information and advice

Sports clothing and equipment trade events
ISPO, Munich, Germany, winter and summer and other global fairs. (www.ispo.com)
European OutDoor Fair, Germany, annual fairs. (www.european-outdoor.de)
Outdoor Retailer, USA. winter and summer fairs. (www.outdoorretailer.com)

Sports related technical textile trade events
Avantex / Techtextil, Frankfurt, Bi-annual.

Journals and publications
Future Materials, World Textile Publications Ltd., Perkin House, 1 Longlands Street, Bradford, West Yorkshire, BD1 2TP, UK.
Textiles Intelligence Ltd., ‘Performance apparel markets: Business and market analysis of worldwide trends in high performance activewear and corporate apparel’. 10 Beech Lane, Wilmslow, Cheshire, SK9 5ER, UK.
Zoom: The Magazine of Schoeller Textile AG, Banhoffstrasse 17, CH-9475 Sevelen

Websites
http://www.schoeller-textiles.com/
http://www.polartec.com/
http://www.baltex.co.uk/
http://www.exo2.co.uk/
http://interactive-wear.de/
16.7 References

17.1 Introduction

It is estimated that about 25% of the total worldwide use of fibre in textiles is used for technical textiles but in advanced countries this figure increases to about 40%. In technical textiles, the largest user is the automotive sector as shown in Table 17.1, whose figures were obtained from statistics in 2003. This sector uses about 2000 kt. The table also shows that the average unit price of fibre for automotive use is comparatively high compared with the main technical-textile end-use fibres. The amount of fibre used for a standard passenger car is about 25 kg. This is now increasing because the requirements for the safety and comfort of passengers and for car weight reduction, which means an increase of reinforcing fibres for hard composites, are being intensified.

Furthermore, the worldwide demand for automobiles is growing, as shown in Fig. 17.1. Hence, the amount of automotive fibre being used is therefore increasing. These statistical facts indicate the industrial importance of automotive fibre in terms of end-use.

The selection criterion for automotive use materials is ‘performance/cost’ as with most other technical textile materials. In general it is more strictly applied to automotive use. For example, steel cord is usually used as a reinforcing material in tyre belt plies for passenger cars. The reason is that steel has the best value for ‘(tensile modulus)/cost’ of currently existing materials. But if car weight reduction becomes more important in the future, the selection criterion may become ‘(specific tensile modulus)/cost’ and then steel could be replaced by a high-performance fibre such as poly-ketone fibre.

Fibre is used for several kinds of parts in automobiles such as (a) tyres, driving belts, tubes and hoses, seat belts, air bags, seats, roof trims, floor coverings, noise control materials and cover sheets, (b) filters, and (c) mechanical parts, exterior body panels and bumper beams. In the near future, optical fibre for the information and driving control system, and
reinforcing fibre for the fuel pressure vessel may also be widely used. The percentage use of automotive fibres for main parts is summarized in Table 17.2. Hard composites is the second largest user group but they are mostly reinforced by glass fibre through compression moulding and/or injection moulding. As described in Section 17.3.1, nylon tyre cord is preferable for use when driving on rough roads, especially in developing countries, because nylon is superior in toughness compared with PET. On the other hand, rayon fibre is preferred for use in run-flat tyres in Europe and North America, because it has better heat resistance than PET. With regard to the shares in the total amount of automotive fibres used, polyethylene

\[ \text{Table 17.1 Share in consumption amount and relative unit price by end-use field in technical textiles}^1 \]

<table>
<thead>
<tr>
<th>End-Use Field</th>
<th>Share (%)</th>
<th>Relative Unit Price</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transportation</td>
<td>25.3</td>
<td>3.5</td>
</tr>
<tr>
<td>Industrial</td>
<td>15.6</td>
<td>2.2</td>
</tr>
<tr>
<td>Sports</td>
<td>15.3</td>
<td>3.8</td>
</tr>
<tr>
<td>Living goods</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>Construction</td>
<td>7</td>
<td>1.3</td>
</tr>
<tr>
<td>Clothing goods</td>
<td>7</td>
<td>1.9</td>
</tr>
<tr>
<td>Sanitary, medical</td>
<td>6</td>
<td>1.2</td>
</tr>
<tr>
<td>Agriculture, fishery</td>
<td>6</td>
<td>1.4</td>
</tr>
<tr>
<td>Protection</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>Packaging</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td>Civil engineering</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

\[ \text{Figure 17.1 Trend in the demand for automobiles in the world}^2 \]
Polyester and polyamide fibres for automotive use in terms of fibre performance requirements

17.2.1 PET, nylon 6 and 66 fibres

In Table 17.3, the main properties of PET, nylon 6 and nylon 66 fibres are shown compared with those of polypropylene (PP) and p-aramid fibres. In Table 17.4, the advantages and disadvantages of PET, nylon and PP fibres in terms of properties, cost, and energy consumption are summarized.

PP fibre is usually the most economical, lowest in energy consumption to produce fibre and lowest in specific gravity. But PET and nylon fibres have advantages in dyeability, high temperature resistance and dimensional stability over PP. It must be noted that the lower melting point of PP causes a limitation in the application of PP to temperature-sensitive parts. Compared with nylon fibre, PET fibre has a higher modulus, higher heat stability, higher resistance to colour change, higher durability for...
### Table 17.3 Properties of PET, nylon fibres comparing with PP and p-aramid fibres (standard type)

<table>
<thead>
<tr>
<th>Fibres</th>
<th>Specific gravity</th>
<th>Melting point (°C)</th>
<th>Glass transition temp (°C)</th>
<th>Tensile strength (MPa)</th>
<th>Breaking elongation (%)</th>
<th>Tensile modulus (GPa)</th>
<th>LOI (%)</th>
<th>UV degradation</th>
<th>Colour-change by UV</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET</td>
<td>1.38</td>
<td>260</td>
<td>70</td>
<td>510–690</td>
<td>15–40</td>
<td>6–11</td>
<td>18–21</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Nylon 66</td>
<td>1.14</td>
<td>260</td>
<td>35 (50%RH)</td>
<td>350–550</td>
<td>18–36</td>
<td>3.0–6.5</td>
<td>20–21</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Nylon 6</td>
<td>1.13</td>
<td>220</td>
<td>20 (50%RH)</td>
<td>450–700</td>
<td>20–32</td>
<td>2.5–3.4</td>
<td>20–21</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>PP</td>
<td>0.91</td>
<td>160</td>
<td>−15</td>
<td>410</td>
<td>25–60</td>
<td>6.4</td>
<td>18–20</td>
<td>+*</td>
<td>+++**</td>
</tr>
<tr>
<td>p-Aramid</td>
<td>1.14</td>
<td>–</td>
<td>300</td>
<td>2760</td>
<td>3.3</td>
<td>58</td>
<td>29</td>
<td>++</td>
<td>−</td>
</tr>
</tbody>
</table>

**Notes:**
++: fairly good; +: good; +*: can be good by additive; +++*: very good by pigment colour; −: sensitive
sunlight degradation and is less expensive. In addition, the production of nylon fibre uses the most energy. But it has greater toughness, excellent tensile strain recovery and also excellent adhesiveness. With regards to nylon 6 and 66, the former has higher heat resistance but it is a little more expensive. Hence, nylon 66 is used for the end-uses requiring higher heat resistance. In these property and cost situations, PET fibre is usually used for the reinforcement of most rubber composite parts. But for tyres used on rough roads in developing countries, advantage is taken of the excellent of its higher modulus, higher glass transition temperature and higher durability for sunlight. On the other hand, nylon 66 fibre is used for air bags because of its higher foldability when it needs to be contained and higher resistance to small burning particles. For car interiors, PET fibre is mainly used because of its higher sunlight (colour change) resistance and higher heat resistance.

### 17.2.2 Other polyester fibres and polyamide fibres

Polyethylene naphthalate (PEN) is the other main type of polyester fibre which is now used in automotive materials. As shown in Fig. 17.2, PEN has a much higher tensile modulus and heat stability than PET because of the higher molecular stiffness caused by the naphthalene ring for PEN compared to the benzene ring for PET. Its cost is situated midway between PET and p-aramid. One example of its automotive applications is in tyre

Table 17.4 Advantages and disadvantages in properties of PET and nylon fibres comparing with PP fibre

<table>
<thead>
<tr>
<th>Fibres</th>
<th>Advantage, feature</th>
<th>Disadvantage</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET</td>
<td>Balanced in physical properties</td>
<td>Easy in hydrosis, A little less in wearing resistance</td>
</tr>
<tr>
<td>Nylon</td>
<td>Excellent in toughness and wearing resistance, Low in modulus, Excellent in strain recovery</td>
<td>Easy in yellowing, Fairly expensive as conventional fibres, High in energy consumption for fibre production</td>
</tr>
<tr>
<td>PP</td>
<td>Excellent in chemical resistance, Low specific gravity, Low in energy consumption for fibre production</td>
<td>Very bad in dyeability, Low in heat resistance, High in creep and stress relaxation</td>
</tr>
</tbody>
</table>
cap plies because its high modulus can reduce the noise caused by the vibration of tyres in contact with the road. The fibre is also useful for the reinforcement of other automotive rubber composite parts such as brake hoses and driving belts. It should be noted that there are wholly aromatic polyesters and p-aramids other than PET, PEN and nylon 6 and 66. But all of them are generally too expensive to be widely used as automotive materials. Some high-class tyres and high-performance tyres use p-aramid fibre. One typical example of its use is as a cap ply in combination with nylon 66.

### 17.3 Rubber composite parts

#### 17.3.1 Tyres

Tyre structures can be classified as bias tyres and radial tyres. These are illustrated in Fig. 17.3. A bias tyre has a higher energy absorption capacity, therefore it is more feasible to use it on rough roads. But it also has a lower wearing resistance. Hence it has been gradually replaced by the radial tyre with the increase in smoother road surfaces. In the bias tyre, nylon fibre is used for body ply cord because of its excellent toughness. In radial tyres for passenger cars, PET fibre is mainly used for body ply cord because its higher modulus can contribute to improved driving comfortability by the reduction of flat spotting. Nylon 66 fibre is mainly used for cap ply cord because of its higher strength and higher toughness. Figure 17.4 shows the changes by year in the amounts of several tyre cord materials used in USA,
which correspond to the historical changes of the bias tyre and radial tyre, as stated above.

A high modulus low shrinkage (HMLS) type of PET multi-filament is usually used for tyre cord, because a higher modulus and higher heat resistance in shrinkage are important requirements for tyre cord. But the most important property that needs to be improved for PET is its comparatively low adhesiveness to matrix rubber. As regards nylon, a high strength type of multi-filament is required for tyre cord. The properties of tyre cords made of several materials are summarized in Table 17.5 in which yarn.

17.3 Structures of tyre for a passenger car: (a) bias tyre; (b) radial tyre; (c) the cross-sectional structure of a radial tyre.
thickness/number of ply is also shown. There are different methods for the fibre reinforcement of tyres other than by using cord. One is a direct blend of short fibres such as p-aramid with matrix rubber. Another example is the use of the composite made of p-aramid nonwoven and rubber which is inserted into the inner part of a truck/bus tyre to increase run-flat capability.

Table 17.5 Properties of nylon 66 and HMLS PET tyre cords comparing with other materials

<table>
<thead>
<tr>
<th></th>
<th>Ny66</th>
<th>HMLS PET</th>
<th>PEN</th>
<th>Rayon super III</th>
<th>p-Aramid</th>
<th>Steel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thickness</td>
<td>1400/2</td>
<td>1670/2</td>
<td>1670/2</td>
<td>1840/2</td>
<td>1670/2</td>
<td>–</td>
</tr>
<tr>
<td>Twist (T/10 cm)</td>
<td>39 × 39</td>
<td>39 × 39</td>
<td>39 × 39</td>
<td>47 × 47</td>
<td>32 × 32</td>
<td>–</td>
</tr>
<tr>
<td>Strength index of flat yarn</td>
<td>100</td>
<td>90</td>
<td>80</td>
<td>55</td>
<td>210</td>
<td>–</td>
</tr>
<tr>
<td>Elongation of flat yarn (%)</td>
<td>19</td>
<td>11</td>
<td>9</td>
<td>9</td>
<td>3.6</td>
<td>–</td>
</tr>
<tr>
<td>Therm. shrink. of flat y. (%)</td>
<td>5.5</td>
<td>3.0</td>
<td>1.5</td>
<td>≥0.3</td>
<td>≥0.3</td>
<td>–</td>
</tr>
<tr>
<td>Elongation at fixed load (%)</td>
<td>7.0</td>
<td>4.0</td>
<td>2.0</td>
<td>2.0</td>
<td>0.9</td>
<td>–</td>
</tr>
<tr>
<td>Creep index</td>
<td>100</td>
<td>50</td>
<td>–</td>
<td>70</td>
<td>10</td>
<td>–</td>
</tr>
<tr>
<td>Adhesiveness</td>
<td>good</td>
<td>fairly g.</td>
<td>fairly g.</td>
<td>good.</td>
<td>fairly g.</td>
<td>good</td>
</tr>
<tr>
<td>Specific gravity</td>
<td>1.14</td>
<td>1.38</td>
<td>1.36</td>
<td>1.52</td>
<td>1.44</td>
<td>7.81</td>
</tr>
<tr>
<td>Melting point (°C)</td>
<td>265</td>
<td>260</td>
<td>272</td>
<td>degrade</td>
<td>degrade</td>
<td>1450</td>
</tr>
<tr>
<td>Glass transition temp. (°C)</td>
<td>–</td>
<td>70</td>
<td>113</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

Notes:
Rayon super III: the third generation super tenacity rayon
17.3.2 Driving belts

Driving belts can be generally classified as V-belts, V-ribbed belts, cogged belts and metal combined belts, which are shown in Fig. 17.5. For subsidiary automotive equipment such as air conditioners, a V-belt is used. Cogged belts and V-ribbed belts are used for engines. Recently a metal combined belt has been used for speed variation in small passenger cars. PET fibre is applied to the cords of V-belts, V-ribbed belts and metal combined belts.

17.4 Internal safety systems

Seat belts and air bags are very important as passive safety systems inside automobiles.
17.4.1 Seat belts

A seat belt is used to prevent a passenger in a car being thrown from their seat in an accident by fixing them to the seat and absorbing the impact shock. Figure 17.6 illustrates an example of a seat belt. PET woven fabrics are usually used for its webbing. As described in Section 17.2.1, PET is more suitable than nylon for the webbing, because PET has a higher impact energy absorbing capability and suffers less discoloration by sunlight. There are several standards for webbing such as the Federal Motor Safety Standard, Economic Commission for Europe and JIS. The main mechanical requirements in these standards are strength, width, elongation and energy absorption ratio. There are also requirements related to durability such as wearing resistance, cold and heat resistance, water resistance, light degradation resistance and colour fastness. Typical examples of specification values of PET fibre for this end use are: yarn thickness 1000–2500 dtex with 4–25 dtex of its mono-filament thickness, and strength 8–10 cN/dtex.

17.4.2 Air bags

The function of an air bag is to cushion a passenger from collision impact. The bag is instantly inflated by heated gas after a collision stronger than a certain limit. An air bag can be very effective as a passive safety system in combination with a seat belt. There are several kinds of air bags: driver bag, front passenger bag, thorax bag, curtain bag, rear bag and knee bag. Figure 17.7 shows photographs of several kinds of air bag. The base ma-
Material of an air bag is nylon 66 weave because nylon is more suitable for use in an air bag than PET as described in Section 17.2.1. The thickness of nylon filament used for air bags ranges from 235 dtex to 700 dtex. The number of mono-filaments ranges from 70 to 220. A high strength type is needed. There are two types of base material: coated and non-coated. The coating material is now mainly silicone resin. The advantages of the coated type are: better non-gas-permeation, easier bag pressure control, greater heat resistance to burning particles. On the other hand, the non-coated...
Polyesters and polyamides

type is lighter in weight, thinner, more flexible and less expensive. For non-coated cloth, some special products have been developed by weaving with an ultra-high yarn density, with a filament of lower mono-filament thickness, and with a filament whose fibre cross-section is flat.

17.5 Car interiors

The main interior components of an automobile are the seats, door trim, roof trim and floor covering.

17.5.1 Seat and door trim

Polyester fibre is used for most of the seat skin sheet, and for some seat cushion material. It is also used for some door trim skin material. There are several kinds of seat skin sheets such as pile weave, weave, tricot with raising, pile double raschel knit, and pile circular knit. General trends are towards an increase in knit fabrics (tricot, double raschel and circular knit) which are less expensive and have more formability than weave fabrics. Recently a suede fabric, using a PET nonwoven base material, has been introduced. Skin sheets containing phase change material have been also been developed, which can increase the temperature comfortability of the seat. The essential properties of seat skin sheets can be classified as aesthetic effect, physiological comfortability, strength/wearing durability, colour fastness, flame retardancy, heat resistance and non-volatile substance content. They are also often required to have some special functions such as anti-bacterial, deodorizable, anti-static and stain resistant properties. Pile or raising of these sheet fabrics is usually related to an increase in the values of tactile and visual aesthetic effects. An extremely high level of colour-fastness in sunlight is usually required, because automobiles can be used in high temperatures and strong sunlight.

Urethane foam is usually used for seat cushion material under the skin sheet. But some special PET nonwovens have recently been developed such as a fibre mass stabilized with elastomer fibre bonding, a folded web and stitch-bonded web, and a PET 3-dimensional knit fabric with super water absorbancy. One of the most important advantages of these fibrous materials over urethane foam is their moisture permeability for passengers’ physiological comfort.

The material of door trim skins is usually made of plastic such as vinyl chloride and polyolefin. But textile sheets are also used for higher-class cars. In most cases, the textile material is the same as the seat skin fabric. However, the fabric needs to have high enough formability to be made into the complicated shape of a door trim. Its lower end is usually covered by the same carpet as the floor because the door often gets kicked.
17.5.2 Roof trims

PET nonwoven and tricot are used as roof trim skin sheets. The use of needle-punched nonwoven in particular has increased, and nonwoven patterned into velour by needling is especially common. But there are also spun-laced nonwovens and stitch-bonded nonwovens. They are usually formed into roof trim by integration with base materials. Pigment-dyed PET has been widely adopted. Sheets need to have colour-fastness for sunlight, heat resistance, mechanical durability, light durability, formability and non-volatile substance content and stain resistance in addition to lightness. Certain levels of sound absorbing capability and heat insulation are also needed. The base materials for roof trim can be mainly classified as polymer foam sheets and fibre-reinforced porous polymer sheets. Glass fibre is used for most of these sheets.

17.5.3 Floor coverings

Floor coverings can be mainly divided into liner coverings and optional coverings. Needle-punched nonwoven carpet and tufted carpet are used for liner coverings. The usage of nonwoven carpet has greatly increased because it is more economical and has better formability. There are three main kinds of surface pattern: plain, loop-like and velour-like, as schematically shown in Fig. 17.8. The latter two patterns can be made by the combination of specific needles and needling patterns. The surface material of these needle-punched carpets mainly consists of pigmented PET fibre and/or pigmented PET recovered from bottles. Its fibre thickness is reduced

![Diagram of three main types of surface patterns for needle punched carpet: (a) plain; (b) loop-like; (c) velour-like.](image-url)
from 11 to 6 dtex to increase its covering factor. Quite attractive aesthetic effects can be achieved with needle-punched carpet by the specific patterning described above. The main performance requirements are: mechanical durability, sunlight durability, heat resistance, sunlight colour fastness, sound absorbability, sound insulation, anti-fogging, non-volatile substance content, flame retardancy, stain resistance and formability. With an increased requirement for weight reduction, sound absorbability has become more important than sound insulation.

Tufted carpet has usually been used for optional carpets, where its aesthetic effect and appearance of high quality to the customer is of greater importance. Its weight ranges from 350 to 2000 g per m² based on the wide variety of customer requirements. Its surface material fibre is nylon, PET or PP. Of these, PP is increasing its usage share.

### 17.6 Others

#### 17.6.1 Exterior cover sheets

There are two kinds of automotive exterior cover sheets. One type is a cover for protecting lorry cargo, which is further subdivided into covers for hood use and for plain covering use. The other type is for protecting passenger cars. PET spun yarn weave is mainly used for lorry cargo covers. Figure 17.9 shows its typical structure. Its coating layer is usually poly-vinyl chloride. The main specifications of cover sheets are also summarized in Table 17.6. Some PET filament weave is also used for truck cargo covers but filament weave is inferior in durability to spun yarn weave. The main performance requirements are: waterproof properties, water repellency, mechanical durability, weather resistance, dimensional stability, stain resistance, ability to be welded by radio frequency, and lightness.

Cover sheets for protecting passenger cars are usually made of PET filament weave, whose yarn thickness is about 250 dtex. Its weight is about 200 g per m². It is usually coated to make it waterproof. Some cover sheets

![Cross-sectional structure of exterior cover sheet](image-url)
Some other cover sheets are flame-retardant to protect cars from fire.

17.6.2 Boot mats

The performance requirements for a boot mat are not great enough to make it a particularly cost-sensitive automotive part. Needle-punched non-woven is usually used for the mat. Felt made of recovered fibre is also used for the underlayer of the mat.

17.6.3 Noise control materials

The sound insulation capability of a sheet is generally proportional to its weight. For example, 10 kg per car is needed for a good sound insulation floor covering. In order to reduce car weight, the strategy for noise reduction within the cabin has been changed to adopting the use of parts having higher sound absorption capability instead of higher sound insulation. Interior parts such as floor covering and roof trim are typical targets in this strategy, as stated above. Multi-layered nonwovens are used for the engine silencer and sheet supporting spring to reduce unusual sounds in line with this strategy. Figure 17.10 is an example of such an engine silencer.

17.7 Conclusion

In the future, the need for eco-friendliness in cars will become more important. Weight reduction will be attempted for all kinds of automotive parts and for car structures. Trials on the structure of polymer material will be carried out for its ability to be recovered easily from scrapped cars.
The re-use of recovered materials for automobiles will be increased. Carbon neutral material such as poly-lactic acid and natural cellulose fibre may be gradually introduced into automotive parts. These factors might have positive impacts in some cases and negative in other cases on the automotive use of polyester fibres and nylon fibres. But it is most likely that the use of these fibres will not be particularly affected, at least in the near future. Therefore, the amounts of these fibres for automotive use will increase at least in proportion to the increase in car demand.

Many of the references in this chapter are unfortunately written only in Japanese. Hence the author would like to list some books written in English at the end of the following references for the convenience of readers.

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18 Applications of polyesters and polyamides in civil engineering

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18.1 Introduction

A niche market for the textile industry is providing fibres and fabrics to the construction industry, as a potential substitute for more traditional building materials such as wood, concrete, masonry and steel. If in several applications this is the main goal of the textile industries, in other cases, textile materials are already considered traditional civil construction materials.

Fibre reinforced composites are gaining popularity within the civil construction sector. Applications like fibre-reinforced concrete, concrete retrofitting, concrete jacketing and composite concrete structures internal and external reinforcement are, among others, applications where fibres and fibrous structures play an important role. In these emerging applications fibres like AR-glass, carbon and aramid are of paramount importance due to their high strength, high modulus and to the possibility to compete with the conventional structural construction materials.

In applications related to the geotechnical and geoenvironmental applications, where properties beside mechanical ones are of paramount importance, fibres and textile structures are already commonly used materials. Examples like embankments and retaining walls, road construction and road overlaying, erosion control of steepened slopes, shore protection, construction of waste landfills, architectural membranes, offshore applications, among others, show that fibres and fibrous structures are the core construction materials. In these applications polyamide fibres and, especially, polyester fibres are used in an indiscriminate way.

18.2 Polyester and polyamide fibres and structures for civil construction applications

Depending on the applications envisaged, polyester and polyamide fibres are used in civil construction projects in several forms. If concrete internal
reinforcement is the objective, polyester and polyamide fibres are used as micro- and macro-fibres, fibrillated fibres and monofilaments. Otherwise, if concrete external reinforcement is required, polyester and polyamide fibres are applied as woven fabric. Polyester and polyamide woven and knitted fabrics as well as needle punched nonwoven fabrics, are also available in geotextiles and geogrids. The combination of different types of structures – hybrid structures – is also common, and their impregnation in matrices, polymeric or cementitious, is usual.

18.2.1 Monofilaments, micro- and macro-fibres and fibrillated fibres

Polyester and polyamide fibres are available in several geometries and dimensions. Monofilaments, rovings (tows) and fibrillated fibres are specially used to reinforce cementitious or polymeric matrices. Monofilaments are single straight fibres usually made by drawing molten polymers. Such fibres are usually round in cross-section and are usually used in fibre-reinforced concrete applications. Many polymers are spun in the form of filaments whose diameters vary from 5 to 15 μm in diameter. Several thousand filaments are spun simultaneously, by drawing them from a single die and bundling together. Rovings are cut into short lengths in order to obtain micro- and macro-fibres. Dispersed rovings act as individual monofilaments. The geometry of a fibrillated fibre can be described in terms of a film thickness and the width of each filament.

18.2.2 Fibrous structures

Polyester and polyamide fibres are available in several fibrous structures. The most used polyester and polyamide fibrous structures are geotextiles and geogrids. Geotextiles can be woven, knitted or nonwoven structures. Geogrids can be produced with conventional weaving and/or knitting process, coated with copolymer resins.

Conventional woven fabrics are produced by orthogonal interlacing two sets of yarns: one called warp whose yarns are along the length of the fabric, while the other one is weft whose threads run along the width of the fabric. Both sets of yarns are mutually positioned under the angle of 90° (warp at 0° and weft at 90°). The warp and weft yarns can be interlaced in diverse ways presenting a regular pattern named woven weave structure. Plain weave is the simplest woven structure and it is characterized by the interlacement of warp and weft yarns, alternately over and under. This is a balanced structure, with good stability and reasonable porosity. However,
it presents several disadvantages such as difficulty to drape and high level of fibre crimp leading to low mechanical properties.

Woven fabrics are usually dimensionally stable, but they present lower extensibility and porosity rather than other structures. The mechanical properties of woven fabrics are very important for technical textiles and they depend on different factors, namely raw materials, linear density of the warp and weft, yarn density and woven structure. The strength of a fabric is usually higher in the warp and weft directions, while in diagonal direction they present lower mechanical properties, higher elasticity and lower shear resistance.

Knitted fabrics are textile structures characterized by the interlacing of loops. Two different basic textile technologies allow the manufacturing of knitted structures: weft and warp knitted technologies. Both types of knits produced by these technologies can have extensibility in one specific direction and by using straight yarns, which are not knitted, it is possible to design the stability in one direction, with deformability in the others [1]. All loops of a warp knitted fabric are formed from a separate yarn, called warp, mainly introduced in the longitudinal fabric direction. Therefore, neighbouring loops of one course are not produced from the same yarn. These structures have mechanical properties very similar to those of woven fabrics, are very flexible and, depending on the construction, can be elastic or inelastic. For the production of weft-knitted fabrics, a single yarn may lead to the formation of an entire course (horizontal row of loops). One important factor that influences the properties of weft knitted fabrics is the stitch length, that is the length of a yarn in a knitted loop. The weft-knitting technology presents great ability to produce complex shapes which is very important for technical applications.

Nonwoven is a very interesting production technique due to the wide variety of products that may be developed comparing to other textiles technologies. According to EDANA [2], nonwovens are sheets, webs or bats of natural and/or man-made fibres or filaments, that have not been converted into yarns, and that are bonded to each other by entangling fibre or filaments mechanically, thermally or chemically [2]. The process of nonwovens manufacture can be divided in two steps: the matt preparation (carding, air laying, wet laying or spun laying) and the bonding process itself. Needle-punched nonwoven geotextiles are entangled to form a complex structure by random fibres, accounting for its bulky nature, wide range of pore size distribution and good drainage.

Braided textile structures are produced by intertwined yarns. There are three types of braided structures: diamond, regular and hercules. Diamond structure is obtained when yarns cross alternately over and under the yarns of opposite direction and, in this case, the rotation is one to one. But it is possible to modify this rotation to two by two and three by three
obtaining, in this way, the other two structures. Among these structures, diamond is the most stable and hercules presents the least stability. Usually, braids are produced in a biaxial structure; however, if yarns are inserted (middle-end-yarn) and oriented longitudinally into the structure, a three axial braid is obtained. Also it is possible to insert in the middle of the tubular form fibre bundles, obtaining in this way a triaxial structure with axial fibres. The diameter of the braid is controlled by several factors namely, the number of fibres, the angle of the intertwining yarns, the number of intersections of fibre per unit length of the yarns and the linear density of the fibres. Braiding technology has been used for the production of hollow pieces with or without core. The braid structures have weak axial and compression stability and are conceived to have multidirectional conformability [1]. Nowadays, braids are the reinforcements with more variety of applications on the market, namely the rope and offshore industries.

18.3 Synthetic fibre-reinforced concrete

The term fibre-reinforced concrete is defined by ACI 116R, Cement and Concrete Terminology, as a concrete containing dispersed randomly orientated fibres (Figure 18.1) [3]. Fibre reinforced concrete can be described as a composite material consisting of a concrete phase and a small portion of discrete and discontinuous fibres distributed and oriented randomly within the concrete [4]. The development of steel fibre reinforced concrete may have begun around the early 1960s. Polymeric fibres came into commercial use in the late 1970s, glass fibres experienced widespread use in the 1980s, and carbon fibres attracted much attention in the 1990s [5, 6].

18.1 Synthetic fibre-reinforced concrete.
Inherently concrete is brittle under tensile loading and mechanical properties of concrete may be improved by randomly oriented short discrete fibres which prevent control initiation, propagation, or coalescence of cracks [3, 7, 8]. The major disadvantage of concrete may be thought to be the complexity of its structure, which results in many internal stress concentration zones [5]. Thus, some internal micro-cracks exist at cement paste-aggregate interfaces even prior to any load and environmental effects. Under externally imposed structural loads and environmental effects, concentration of tensile stresses occurs at the interfaces between aggregates and matrix, causing the growth of micro-cracks in size and number; propagation of interface micro-cracks into the matrix and eventual joining of micro-cracks yield large cracks and lead to the failure of the concrete. Cracks in concrete structures reduce safety and durability even during their life time [9]. After the formation of the first crack, brittle matrices, as plain mortar and concrete, lose their tensile load-carrying capacity almost immediately [10]. The cracks generally develop with time, thereby impairing the waterproofing properties and exposing the interior of the concrete to the destructive substances containing moisture, bromine and acid sulphate, among others. The exposure acts to deteriorate the concrete, with the reinforcing steel corrosion [11, 12].

Fibre-reinforcement is used to improve the brittle nature of cementitious composites [9]. Fibre-reinforced concrete is more ductile than unreinforced concrete. One way to quantify the increase of ductility is to measure the area under the load-deflection curve, which is defined as toughness. Fibres bridge cracks during loading and transfer the load, arresting the growth and coalescence of cracks. Fibre reinforcement has been shown to improve the ductility, toughness, flexural strength and shear strength of cementitious materials, to reduce shrinkage cracking and permeability and enhance fatigue and impact resistance. Fibres are added not only to improve tensile strength [5, 13]; recently, it has been found that a number of fibres can also improve the residual properties of concrete after exposure to elevated temperatures [14].

Synthetic fibres have become more attractive in recent years as reinforcements for cementitious materials. Synthetic fibres can provide effective, relatively inexpensive reinforcement for concrete and are alternatives to asbestos, steel and glass fibres. Recently, structurally efficient fibres have been developed, exhibiting increased toughness and/or loading carrying capacity after cracking [15, 16]. Fibre types that have been incorporated into cement matrices include polyethylene (PE), polypropylene (PP), acrylics (PAN), poly(vinyl alcohol) (PVA), aramid, carbon, polyamides (PA) and polyester (PES) [17]. The properties of synthetic fibres vary widely, in particular with respect to the modulus of elasticity, an important characteristic when fibres are used for producing composites. Besides physical
and mechanical properties, durability of fibres used for fibre-reinforced concrete is also of paramount importance [6, 17].

To increase the strength of their composites, fibres must have a modulus of elasticity greater than the matrix. For cementitious materials, where the modulus of elasticity ranges from 15 to 30 GPa, this condition is difficult to meet with most synthetic fibres. Therefore, attempts have been made to develop fibres with a very high modulus of elasticity for cement reinforcement. However, both theoretical and applied research have indicated that, even with low modulus fibres, considerable improvements can be obtained with respect to the strain capacity, toughness, impact resistance and crack control of the fibre-reinforced concrete composites. In many applications, the enhancement of these properties is of much greater significance than the modest increase in tensile (flexural) strength. In some applications, the poor tensile strength of concrete can be enhanced by combining concrete with small diameter fibres, which can be synthetic fibres. It is usually assumed that the fibres do not influence the tensile strength of the matrix, and that only after the matrix has cracked do the fibres contribute by bridging the crack. Most of the current applications with fibre-reinforced concrete involve the use of fibres ranging around 1% by volume with respect to concrete. More recently, it has become possible to incorporate relatively large volumes (ranging up to 15%) of synthetic fibres into concrete. With such large volumes of fibres it has been noted that the fibres may substantially increase the tensile strength of the matrix.

### 18.3.1 Type of reinforcement

Synthetic fibres are usually used as primary reinforcement in thin sheet elements where conventional steel reinforcement is not feasible. These elements are prefabricated rather than cast-in-place. Most structural reinforcements have involved the use of steel, glass or carbon fibres. The development of high modulus synthetic fibres, such as polyaramid, allows the possibility of replacing steel or glass fibres, but at a high cost that makes current use of synthetic fibres for primary reinforcement an unfeasible alternative.

Secondary reinforcement is used to control cracking caused by intrinsic tensile stresses such as drying shrinkage, plastic shrinkage or temperature changes. The purpose of crack control is neither to eliminate cracking caused by intrinsic stresses, nor to increase the load-carrying capacity of the concrete; crack control replaces a random pattern of relatively large cracks with a more deliberately structured pattern of closely spaced fine cracks. Actually, this makes concrete less water permeable and hence more durable.
18.3.2 Effect of fibres on fresh and hardened concrete

Dispersion of fibres during mixing is an important concern in the production of fibre-reinforced concrete. Despite the benefits of fibre reinforcement, fibres can make cementitious materials difficult to work, in the fresh state, compromising hardened state properties [13]. The addition of fibres at high dosages, however, has potential disadvantages in terms of poor workability and increased cost [18, 19]. Micro-defects, such as voids, honeycombs and clumping or ‘balling’ of fibres can be a problem at high fibre contents [6]. A compromise to obtain good fresh concrete properties and good toughness of hardened concrete can be obtained by adding two different types of fibres, which can work individually at different scales to yield optimum performance and can reduce the overall cost of concrete production. The addition of non-metallic fibres such as glass, polyester, polypropylene, etc. results in good fresh concrete properties and reduced early age cracking. It is also important to have a combination of low and high modulus fibres to arrest the micro- and macro-crack, respectively. Another beneficial combination of fibres is that long and short fibres can control different scales of cracking [18, 19]. The beneficial effects of non-metallic fibres could be attributed to their high aspect ratios and increased fibre availability at a given volume fraction. Because of their low stiffness, these fibres are particularly effective in controlling the propagation of micro-cracks in the plastic stage of concrete. However, their contribution to post-cracking behaviour, unlike steel fibres, is not known to be significant [18, 19].

Synthetic fibres seem to perform better than glass or steel fibres. Plastic shrinkage cracking is caused by excessive loss of water from fresh concrete due to evaporation. The problem is most acute in the summer months when weather conditions may increase the rate of moisture evaporation so that it exceeds the rate of bleeding [19]. When it happens, local drying at the surface will result in plastic shrinkage cracking. Once the magnitude of tensile stresses needed to cause cracking is quite low, synthetic fibres have the potential to prevent his type of cracking in cases where inadequate preventive measures have been taken. Laboratory studies show that slabs with fibre reinforcement develop less plastic shrinkage cracking than slabs of control concrete [6].

Some studies have shown no significant improvements in tensile or compressive strength of concrete when low volumes of synthetic fibres are added to concrete. Other studies have shown a slight decrease in compressive strength, over unreinforced concrete control specimens. These apparent differences are attributed to changes in the water-to-cement ratio between control and fibre-reinforced concrete. Since the addition of fibres slightly decreases slump, additional water is often used to bring the slump
back into the desired range. This small addition is sufficient to lower strengths by about 10% [6].

Several studies reported greater static flexural strength in concrete with added synthetic fibres than in unreinforced concrete. However, these increases were quite small when fibre contents were low. The static flexural strength of fibre-reinforced concrete increases after being subjected to fatigue loading, behaviour also observed in unreinforced concrete [17].

Fibre-reinforced concrete is expected to have a finer crack pattern than plain concrete. The crack spacing is directly related to the radius of the fibre for circular cross-section. The finer the fibre, the smaller the crack spacing. Another factor of paramount importance is the fibre-concrete interfacial bond strength. The reduction in crack width as the crack spacing is reduced will give fibre-reinforced concrete lower water permeability than unreinforced concrete.

18.3.3 Properties of synthetic fibres

Several different kinds of polymers have been investigated for their suitability to use in concrete. In fibre properties evaluation, four aspects are of interest, namely, fibre geometry and dimension, mechanical properties, physical properties and chemical durability.

Fibre geometry and dimensions

A synthetic fibre can be described as a flexible, macroscopically homogeneous body, with a high length to thickness ratio (aspect ratio) and a small cross-section. All fibres have one particular structural feature in common: a preferential orientation of their elemental units with respect to the fibre axis [6].

Fibres are available in many different geometries and dimensions, such as monofilaments, rovings (tows) and fibrillated polymers. Monofilaments are single straight fibres usually made by drawing molten polymers. Such fibres are usually round in cross-section, ranging in diameter from 50 μm to 0.5 mm, being usually strong. However the smooth surface may result in a low interfacial bond strength, which limits the reinforcing capability of the fibre. Bond strength is primarily determined by nature of the polymer. If bond strength at the interface between fibre and matrix is too high, fibre ruptures after the first crack initiates. On the other hand, fibre is easily pulled out if the bond strength is too low. Bond strength may dominate the mechanical properties of fibre-reinforced concrete in this way and the combination of fibre and matrix needs to be selected prudently to get efficient bond strength [9]. Rovings of polyaramid present the highest bond strength, slightly higher than polyacrylic monofilaments. Polyamide and
polyester monofilaments present one of the lowest bond strengths (Figure 18.2) [6].

Many polymers are spun in the form of filaments (fibrils) which vary from 5 to 15 μm in diameter. Several thousand fibrils are spun simultaneously, by drawing them from a single die and bundling together. Rovings are cut into short lengths when adding them to concrete. In the concrete, the yarn separates into individual fibrils which need to be dispersed efficiently during mixing. Dispersed rovings act as individual monofilaments within the cement matrix (Figure 18.3). The high surface-to-volume
ratio should increase the effective interfacial bonding. The geometry of a fibrillated fibre can be described in terms of a film thickness (generally 15 to 100 μm) and the width of each fibril (100 to 600 μm). During mixing, the bundles are opened up by the aggregate to act as individual fibrils contributing, each one, to the reinforcing action (Figure 18.4). The fibres are easy to handling and to disperse uniformly within the concrete mix.

**Mechanical properties**

The fibre mechanical properties determine its potential as a concrete-reinforcing material. Commodity synthetic fibres are characterized by low modulus of elasticity and high elongation. Since the modulus of elasticity for most cementitious materials ranges from 15 to 30 GPa, the relatively low modulus of the fibres means that high strength composites are not achievable with synthetic fibres. Their advantage lies in increasing such properties as strain capacity, toughness, and crack control, properties that are more important for some applications. Polyamide fibres present a tensile strength of 1000 MPa, modulus of elasticity of 6 GPa and ultimate elongation of 10% (Figures 18.5, 18.6 and 18.7) [6]. Polyester fibres present a tensile strength of 1100 MPa, modulus of elasticity of 10 GPa and ultimate elongation of 24% (Figures 18.5, 18.6 and 18.7) [6].
Polyesters and polyamides

Physical properties

Fibres are characterized by low density so that a relatively low mass of fibres yields a high volume of fibres in concrete (Figure 18.8). The density of polyamide and polyester fibres are 1.14 and 1.35 g/cm³, respectively [6].

Synthetic fibres are also very flexible; fibre breakage or mechanical distortion will not be a problem during concrete mixing. The relatively low
thermal stability of polymers could be a disadvantage. Melting points are generally below 300°C but service temperatures are considerably lower, since fibres start to soften and lose their tensile properties at temperatures considerably below melting point. This behaviour causes loss of reinforcing capabilities in structures exposed to high temperatures (e.g. fire), but in
Polyesters and polyamides

The glass transition temperature ($T_g$) which represents a change in consistency from rubbery to brittle as the temperature is lowered, should also be considered. Polyamide and polyester glass transition temperatures are 45 and 70˚C respectively [6] (Figure 18.9). In most cases, glass transition lies just above maximum ambient temperature usually encountered in service, as in the case of polyamide, solar heating could cause the temperature in the concrete to exceed $T_g$. Maximum service temperature of polyamide and polyester fibres is 120 and 150˚C respectively [6] (Figure 18.10).

**Environmental durability and thermal resistance**

Cement paste develops a moist alkaline environment detrimental to many organic materials. The minimum pH in a cement paste is about 12.3, which corresponds to a saturated calcium hydroxide solution. However, many types of cement have high alkali content, which can raise the pH to 13.5 or higher, when the cement is mixed with water. This very high pH will be detrimental to some polymers. Polycrylics, polyesters and polyamides are particularly sensitive since they can undergo alkaline hydrolysis [6]. Hydrolysis may be slow at room temperature, but may be significantly accelerated at higher temperatures. Moreover, how much this phenomenon occurs depends on three factors: the degree of saturation of the concrete; the rate of hydrolysis; and the extent that hydrolysis will reduce the mechanical properties of the fibre. Synthetic fibres cannot withstand normal environmental temperature fluctuations will not cause a problem.
High temperatures. Most plastics soften and lose much of their tensile properties at temperatures in the range of 150 to 250°C [6].

18.3.4 Polyamide fibres

Polyamides (PA) are well known as nyons, and are defined as polymers which contain amide groups in the main chain. The commercial success of nyons 6 and nylon 6,6 is due to the outstanding properties of the polymers and an economical raw material base. Commercially, polyamide fibres are available in various lengths in single-filament form. These fibres exhibit low bonding with a cement matrix. Because polyamide fibre has a relatively high cost compared with polypropylene, its commercial potential may be somewhat limited.

Walton and Majumdar [17] demonstrated that the use of these fibres in small amounts substantially improves impact resistance of the composite but has very little effect on its tensile or flexural strength [11]. The high impact resistance of the composite containing nylon fibre is derived from the stretching and pulling-out of the fibres which occurs at large strains after the failure of the matrix and at a lower load. Balaguru [18] compared the load-deformation behaviour of rapid-hardening concrete reinforced with nylon 6 and steel fibres, and found that the post-peak drop is steeper for nylon than for steel fibre.

Song et al. analysed the strength properties of polyamide fibre reinforced concretes [11]. Compared to polypropylene reinforced concrete, the
polyamide fibres claimed a slightly increased ability to distribute themselves throughout the concrete, thus distributing the unfavourable stresses within a greater volume of concrete and improving the concrete’s properties in the plastic and hardened state. The compressive strength of the polyamide reinforced concrete improved by 12.4% over the unreinforced control concrete. Polypropylene reinforced concrete improved the compressive strength by 5.8%. The polyamide- and polypropylene reinforced concretes were 17.1% and 9.7% higher, respectively, than that of the unreinforced control concrete. The strength of the polyamide reinforced concrete posted a 5.9% increase, while polypropylene reinforced concrete registered 1.5%. The polyamide fibres performed better in reducing the incubating of early plastic shrinkage cracks because of their higher tensile strength resulting in a better load transfer across the cracks. Moreover the strength improvements achieved by the polyamide fibres overcome those achieved by polypropylene fibres.

Kurtz and Balaguru [18] studied the postcrack creep of synthetic fibre-reinforced concrete in flexure, comparing the behaviour of polypropylene fibrillated fibres and polyamide single filament (nylon 6). They concluded that both polypropylene and polyamide reinforced concrete can sustain only a small percentage of the postcrack strength. The maximum sustainable stress for the polypropylene represents only 24.9% of the average residual strength and for polyamide reinforced concrete this percentage is 38.3%. It was also concluded that polyamide reinforced concrete was found to creep considerably faster than the polypropylene reinforced concrete but for less time.

18.3.5 Polyester fibres

Polyesters (PE) are defined as polymers containing –CO–O– groups in the main chain. This definition excludes polymers of esters such as vinyl acetate and methyl methacrylate since in these polymers the ester groups reside in the side-chains and not in the polymer backbone. The physical and chemical properties of polyester fibres can be changed substantially by altering manufacturing techniques. Polyester fibres which have a higher modulus of elasticity can provide improvements in impact strength. Polyester fibres provide a higher strength at the beginning of ageing, but the values slightly decrease or remain about the same with accelerated ageing. Patel et al. studied the properties of polyester fibre-reinforced concrete and concluded that the addition of polyester fibre at 1% (by volume) of concrete increased the impact strength by 75%, the split tensile strength by 9%, the flexural strength by 7%, and the compressive strength by 5% [17]. There was no change in modulus of elasticity and shear strength. Polyester fibres are generally not resistant to strong alkalis. Wang et al. [17] studied
polyester fibre in Portland cement concrete and observed that polyester fibres lose strength rapidly in the cement matrix. Therefore, polyester fibres are ineffective without surface treatment as reinforcement in Portland cement products due to their chemical degradation.

There have been several studies [18, 19, 20, 21] on the mechanical properties of various fibre reinforced systems, containing hybrid combinations of steel and non-metallic fibres (glass, polyester and polypropylene) and the plastic shrinkage cracking. Steel reinforced concrete and hybrid reinforced concrete present higher compressive strength than unreinforced control concrete. However, concretes with individual non-metallic fibres do not register any increase in strength compared to unreinforced control concrete. The difference in performance of the steel fibre-reinforced concrete and the hybrid fibre-reinforced concretes, with respect to compressive strength and modulus of elasticity, is not significant. Split tensile strengths of hybrid fibre concretes were found to be higher compared with unreinforced concrete control specimens and monosteel fibre-reinforced concrete. Compared with control concrete, all fibre-reinforced concretes showed a significant increase in flexural strength. The combination of steel and polyester showed the maximum flexural strength. All the fibre-reinforced concretes yield a higher flexural toughness compared with control concrete. The maximum crack width was observed in the case of plain control concrete. Compared with steel reinforced concrete, hybrid reinforced concrete showed better crack control features. The total crack area reduces with the increased addition of non-metallic fibres, where polyester and polypropylene fibres performed better than glass. Although the increase in non-metallic fibres results in a significant crack width reduction, it also causes a decrease in concrete workability. This negative effect restricts the maximum dosage of polypropylene and polyester fibres.

18.3.6 Applications

Polyester fibres are used in fibre-reinforced concrete for industrial and warehouse floors, pavements and overlays and pre-cast products [22, 23]. Polyester micro- and macro-fibres are used in concrete to provide superior resistance to the formation of plastic shrinkage cracks versus welded wire fabric and to enhance toughness and the ability to deliver structural capacity when properly designed, respectively. Polyester macro-fibres can be used as a true alternative to welded wire fabric, steel fibres and conventional light gauge steel reinforcing for pre-cast slabs on grad and shotcrete applications. Monofilament polyester fibres are used as secondary reinforcement thus controlling plastic shrinkage, reducing segregation, minimizing bleeding water, providing a three-dimensional reinforcement versus
two-dimensional with welded wire fabric, reducing cracking, increasing the surface durability and reducing permeability [24]. The primary applications of monofilament polyester fibres are plain concrete (unreinforced concrete), in place of welded wire fabric reinforcement, flatwork, footings, foundations, walls and tanks, concrete pipes, burial vaults and pre-stressed beams.

18.4 Geotechnical and geoenvironmental applications

18.4.1 Geotechnical applications

Geosynthetics are thin polymeric materials that are widely used in civil engineering applications. There are numerous types of geosynthetics available for the design engineer to address the diverse range of applications, and each has a specific function. Functions can be hydraulic (drainage, filtration and waterproofing) or mechanical (protection, reinforcement and separation) [26, 27]. Geosynthetics are being increasingly used as reinforcement in engineering projects such as earth embankments, reinforced soil walls and roadways, among others [28]. Geotextiles are a core member of the geosynthetics family and may be applied as a woven, knitted or nonwoven structure [29].

Among the different geosynthetics products, geotextiles present the widest range of properties. They can be used to fulfil most of the functions, except waterproofing. The use of nonwoven geotextiles for filtration or drainage purposes instead of coarse-grained soils is very attractive in applications such as paved and unpaved roads, landfill covers and lines, earth dams, embankments and retaining walls because of the relative ease of placement and gain in space [27].

Geogrids are classified into three types according to the manufacturing method and form: plastic (sheet extruded from olefin and punched and drawn); textile (high-tensile-strength yarn woven or knitted and then coated by resin) (Figure 18.11); and melting [30].

Mechanically stabilized earth walls: embankments and retaining walls

Mechanically stabilized earth walls have gained popularity over recent decades [32]. Embankments (Figure 18.12) and retaining walls (Figure 18.13) reinforced with geogrids and geotextiles are covered under this category [33]. Geotextiles have been extensively used as drainage and filter materials for the past 40 years. Although the designs of these applications have been based on empirical criteria, they have been overwhelmingly successful [34]. Physical and hydraulic properties of geotextiles are of utmost importance for the design of drains and filters [34]. The filtration
and drainage functions are especially attractive in the construction of soil structures such as retaining walls or embankments. Moreover, geosynthetics materials have strengthened their reputation as reliable reinforcing elements based on improved material characteristics and design practices [35].
There has been an increase of interest in the construction of reinforced soil retaining walls to supporting bridge abutments in place of traditional pile foundations. Two of the main reasons for the increased interest are the reduction in overall cost of the project and the reduction, or potential elimination, of ‘bridge bumps’ which arises from differential settlement between a traditional pile supported abutment and the approach rod [35]. Skinner et al. have studied the design and behaviour of geosynthetic reinforced retaining wall and bridge abutments on yielding foundations [36].

The beneficial effect of geosynthetic material is largely dependent on the form in which it is used as reinforcement. Same geosynthetic material when used in planar layers or discrete fibres will give different strength improvements in different forms. Horizontal geosynthetic layers improve the strength mainly by friction and interlocking between soil and the reinforcement, whereas the randomly oriented fibres improve strength friction and coiling around soil particles [37]. Latha et al. studied relative efficiency of these two forms of reinforcement, i.e. planar and discrete fibre, in improving the shear strength of sand [37].

Two important characteristics of the geotextile must be considered as they relate to their applications: one is the ability to percolate water through the geotextile, generally expressed by the fabric permeability coefficient; the other is the particle retention capacity, expressed by geometrical ratios between characteristic particle dimensions of the soil to be retained and a characteristic opening of the fabric [38]. The particle retention capacity of geotextile has direct relation with its structure, features of the soil to be preserved, as well as conditions of the flowing water in site. The development of the geotextile water retention curves is of great importance to model the transient water flow in earthen systems containing geotextiles where unsaturated conditions may prevail such as in reinforced embankments and retaining walls [39].

Also textile geogrids, made with high tenacity polyester filament, using a conventional weaving and/or knitting process, coated with copolymer resins, are commonly used as reinforcement material in embankments, slopes and retaining walls, among other structures [40]. Typical properties of a polyester geogrid for road, embankments and riverside stabilization applications are presented in Table 18.1 [41].

The required service lifetime of geogrids used for reinforcement of embankment and slopes varies according to the sensitivity of the environmental conditions and required safety factor. Typical lifetimes for slopes and embankments are 50–100 years. Usually, the lifetimes of geogrids are assessed as the long-term creep behaviour which causes shape deformation and collapse of the slopes and embankments [40]. Palmeira et al. studied the drainage and filtration properties of polyester nonwoven geotextiles under confinement. He has concluded that polyester geotextile permittivity
Table 18.1 Technical data of polyester geogrid [41]

<table>
<thead>
<tr>
<th>Properties</th>
<th>Unit</th>
<th>ZHGQ 60/30A</th>
<th>ZHGQ 80/30A</th>
<th>ZHGQ 100/30A</th>
<th>ZHGQ 150/30A</th>
<th>ZHGQ 60/30B</th>
<th>ZHGQ 80/30B</th>
<th>ZHGQ 100/30B</th>
<th>ZHGQ 150/30B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Material</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>High tenacity polyester yarn</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>High tenacity polyester yarn</td>
<td></td>
</tr>
<tr>
<td>Coating</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>PVC coating</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>PVC coating</td>
<td></td>
</tr>
<tr>
<td>Process</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>Knitting</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>Knitting</td>
<td></td>
</tr>
<tr>
<td>Tensile strength</td>
<td>MD</td>
<td>kN/m        ≥60</td>
<td>≥80</td>
<td>≥100</td>
<td>≥150</td>
<td>≥60</td>
<td>≥80</td>
<td>≥100</td>
<td>≥1500</td>
</tr>
<tr>
<td></td>
<td>CD</td>
<td>kN/m        ≥30</td>
<td>≥30</td>
<td>≥30</td>
<td>≥30</td>
<td>≥30</td>
<td>≥30</td>
<td>≥30</td>
<td>≥30</td>
</tr>
<tr>
<td>Extension</td>
<td>MD</td>
<td>%           ≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
</tr>
<tr>
<td></td>
<td>CD</td>
<td>%           ≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
<td>≤13</td>
</tr>
<tr>
<td>Length</td>
<td>m</td>
<td>≤100</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Width</td>
<td>m</td>
<td>≤5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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and transmissivity depend on the geotextile thickness [34]. Jeon et al. also analysed the drainage performance of geotextile composites under confined loads [42]. According to Jeon et al., textile geogrids made from polyethylene terephthalate (PET) high performance yarn coated by resin (polyvinyl chloride or acrylic) are recognized as an important synthetic construction material because PET has high tensile strength and excellent creep properties although it has weak chemical stability. Jeon et al. studied the stability to chemical conditions in the application of textile geogrids to soil retaining wall systems [42].

Road construction, reinforcement and overlaying

Roads and highways are of the utmost importance to the development of any country. Due to systematic traffic of heavy vehicles, climate conditions and mechanical properties of the materials in their constructions, road and highway pavements may last considerably less than expected [43]. The design philosophy of a pavement system depends fundamentally on layers of materials, its rigidity, thickness and resistance to stresses [44].

A good pavement must provide a smooth riding profile, withstand large traffic volumes and transmit the stresses efficiently to the underlying subgrade support. But after a certain number of years, pavement defects (cracking) appear at the surface due to repeated traffic loading, local environmental distress and ageing. The traditional flexible pavement rehabilitation using the overlay method is expensive and rarely provides a durable solution as the cracks rapidly propagate through the new asphalt layer forming the so-called ‘reflective cracks’ (Figure 18.14). Flexible pavements are constructed of bituminous and granular materials. But advent of modern synthetic polymers of increasing thermal resistance has created a growing interest on their applications on pavements. Geosynthetics have been used for unpaved roads on subgrade to fulfil one or more of the basic functions: reinforcement, separation, filtration and drainage [44]. Geosynthetics can be effectively used to: reduce or avoid reflective cracking (Figure 18.14); work as a barrier to avoid pumping of soil fines (Figure 18.15); reduce the pavement thickness by reducing the asphalt cap thickness (Figure 18.16); and increase the lifetime of the pavement.
Geotextiles have been successfully used for reinforcement of unpaved roads of soft subgrade to improve the performance of a reinforced fill layer placed on soft ground [45]. The inclusion of geotextiles and geogrids has the potential to extend pavement life by reinforcing and inhibiting reflective crack-up. The high tensile strength of a reinforced pavement will have significant influence on the behaviour of the pavement surface under traffic loading and environmental movements. Gurung [44] concluded that the use of geosynthetic inclusions increases the tensile strength of the pavement and, when comparing the use of geotextiles and geogrids, he concluded that the tensile peak strength of a pavement reinforced with a geogrid was higher than with a geotextile.

18.4.2 Geoenvironmental applications

_Erosion control of steepened slopes_

Soil erosion along degraded hill slopes has been a problem of serious concern throughout the world. In India, about 5330 million tonnes of soil are being lost every year [46]. Failure of natural slopes is mostly associated with human intervention either for urbanization or for other development activities. Denudation of slopes frequently leads to serious problems of downstream sediment. Denuded slopes are often difficult to revegetate and may become unsightly. Better erosion control methods are needed for construction site slopes, so that the soil movement after shaping will be
minimized and subsequent reshaping will be unnecessary [47]. Unscientific measures of slope protection and improper design of structures will always have a consequence on the stability of slopes. A small initial movement in an unstable slope can trigger further soil water movement resulting in soil erosion and consequences landslides.

A dominant factor affecting runoff and erosion of bare soils is the seal formation on the soil surface due to the disintegration of soil aggregates at the soil surface by impacting drops and the compaction of the disintegrated particles into a thin and very compact sealing layer. To control this process, protection of the soil surface against impacting drops is required, as well as the reinforcement of aggregates at the soil surface. M. Agassi [47] studied the stabilization of steep slopes with geomembranes. Geomembranes are used extensively by soil engineers, e.g., as an interface between compacted and uncompacted soil layers. Geotextile membranes placed to cover the soil surface can absorb raindrop impacts and so control crust formation and reduce the consequent runoff and erosion.

Several materials may be used in steep slopes denuded by erosion to reduce surface soil erosion, but usually they will not yield the vegetation growth. Natural vegetation for sustainable erosion control and slope protection is a proven choice of soil conservation along hilly terrain [46]. The use of geotextiles in steep slopes serves the purpose of protecting the soil and the seeds in the initial stage of vegetation growth [46].

Natural geotextiles made of coir, jute, etc. are preferred to synthetic fibres considering the fact that the material is environmentally friendly and ecologically compatible as it gets degraded with the soil. In an age of growing environmental awareness, synthetic geotextiles present some disadvantages [46].

**Geosynthetics for shore protection**

In recent years, traditional forms of river and coastal structures have become very expensive to build and maintain, because of the shortage of natural rock. As a consequence, the materials used in hydraulic and coastal structures are changing from traditional materials and systems to cheaper ones [48]. Geosynthetic structures for shore protection have demonstrably lower construction and lifetime costs than those of hard structures. Therefore, alternative shore protection structures, especially for sandy coasts, that include geotextile solution are of interest to replace traditional materials and systems, such those constructed from rock or concrete blocks [49].

Recently, woven geotextiles in the form of tubes and bags have been successfully used in various environmental protection facilities, such as underwater breakwater, river dyke improvement, stream bank erosion
control and erosion control of seashores [50]. One of these applications is the use of geotextile tube technology. Geotextile tube technology is mainly used for flood and water control, but it is also used to prevent beach erosion, and for shore protection and environmental applications. Woven, nonwoven and composite synthetic fabrics, i.e. geotextiles, have been used for various types of containers, such as small hand-filled sandbags, three-dimensional fabric forms for concrete paste, large soil, and aggregate-filled geotextile gabion, prefabricate hydraulically filled containers, and other innovative systems involving containment of soils using geotextiles [48].

One innovative system is a geotextile wrap-around revetment which is a sand slope that is reinforced with a geosynthetic: the material is wrapped and encapsulated with geotextiles to create a flexible revetment. A layer of geotextile is first placed at the site and sand is placed on top of it to produce this system. Loose ends of the geotextile are then folded back and inserted into the fill. A second geotextile layer is laid on top and the procedure is repeated until the layers reach the designed height. The finished system is covered with sand to give it a natural appearance. The geotextile encapsulates the sand and adds tensile strength to it through increased confinement [49].

Waste landfills

Geosynthetics have been increasingly used in environmental protection works and in the case of waste disposal areas their functions range from barriers to fluids or gases to reinforcement of earth works and slopes [51]. When disposed in excavations, the volume of waste to be stored can be maximized with the use of steeper slopes, however keeping the appropriate safety margins required in this type of work. The use of geosynthetics in such slopes, as barriers or drainage materials, poses important considerations regarding the stability conditions of the cover soil and adherence between soil and geosynthetics and between different types of geosynthetics [51]. Geosynthetics are modern engineering materials that can be used in this type of problem to work as barriers or in liquid collection systems [52]. They have been increasingly used in liquid collection systems. They are easy and quick to install and, being manufactured, they can be subjected to rigid quality control to guarantee their properties. Among the geosynthetics types, a geotextile can function as drainage or filter layer in a liquid collection system. To perform well as a filter, prolonging the service life of the drainage materials, the geotextile layer has to retain particles from the waste, while maintaining a sufficient permeability to allow the passage of fluid during the entire contaminating lifespan of the landfill [52]. Moreover, besides drainage and filtration nonwoven geotextiles are
widely used in waste landfills as materials having the functions of protection and separation, among others. In general polyester or polypropylene geotextiles are the most important geosynthetic materials that are installed above the geomembranes for protection and drainage. These geotextiles are exposed to chemicals such as acidic or alkaline solutions, especially leachate solutions, until the reclamation of waste is completed [53]. Leachate is a very complex and variable liquid and can bring solid particles in suspension; microbiological activities are intense, biological and physical clogging can occur and leachate flow can be intermittent for most of the disposal area lifetime [52].

Gas drainage layers comprising either natural materials or geocomposites are specified in most modern and recent landfills. Because of the large gas fluxes generated in these landfills, the use of geocomposites or natural material is well indicated to relieve gas pressure underneath the hydraulic barrier in the cover system and allow the collection of gas and its transmission to point outlets. However, for abandoned or uncontrolled old landfill sites, which in most cases generate very low quantities of gas, a nonwoven geotextile can be used as gas collection layer below the hydraulic barrier. Thiel indicated that limited testing of nonwoven needle punched geotextiles showed that these materials may be acceptable for gas relief [54]. A nonwoven geotextile, in general, has relatively large pore sizes compared to most soils, and consequently its gas transmissivity behaviour may be expected to be similar to a coarse soil. Furthermore, its low capillary rise potential, high breakthrough head for saturation and low water retention capacity make it more attractive than conventional coarse-grained soils. Gains in space, cost and construction time are other advantages [54]. Jeon studied eight types of polypropylene and polyester nonwoven geotextiles/geotextile composites.

### 18.5 Textile architectural applications

Architectural engineering is one of the many areas of civil engineering. It is the engineering discipline responsible for the planning, analysis, design, construction, maintenance and renovation of building systems considering always their impact on the surrounding environment [55, 56].

Textile architecture is not a new branch of architecture. Its technological potential is continuously leading to new developments and applications [57]. In order to produce new materials for architectural applications, the research leads to the development of production techniques as well as of opportunities for material applications. Nowadays, designers and engineers are facing an incredible demand for higher efficiency, not only in building construction, but also by the relationship between the construction and the environment [58].
Several textile materials can be used in this area. The continuous development of synthetic fibres allowed the development and the growing of high performance fabrics with excellent properties, suitable for architectural applications, namely, good mechanical resistance, rot and fungus resistance, hydrophobic behaviour, among others [59].

18.5.1 Textile architectural membranes

Since the 1950s, membrane development for building purposes led to new materials with considerable strength and resistance [60]. Textile architectural membranes or tensile membrane structures are normally a woven structure based material, which are often coated or laminated on both sides, in order to improve the mechanical and/or environmental resistance. The main purpose of the coating is to confer a higher durability and impermeability to the fabric and to maintain together all the constituent parts of the membrane. During the coating process, the adhesion stage is extremely important because it will be the indicator of the seam strength. A good adhesion will result in a woven fabric able to carry significant loads. There are several types of coatings. The most common used are polyvinyl chloride (PVC), polytetrafluoroethylene (PTFE or Teflon as commercially known) and silicones. However, for special applications, coating is not applied, but a foil is laminated into the fabric. Lamination process consists in the application of a vinyl film on the woven or knitted structure. Most fabrics used in architecture have a topcoating applied to their exterior coating. The main function of the topcoating is to improve the cleanability of the membrane. Acrylic, polyurethaneacrylic, polyvinyl fluoride (PVF) film lamination or polyvinylidene (PVDF) coatings, are examples of topcoats [59, 60, 61, 62].

In textile architecture, three types of membrane can be used: films, nets and fabrics. Films are polymers in sheet structure without any coating or laminating, including vinyl, polyester and polyethylene. When compared to fabrics, films are cheaper but have low mechanical resistance. Nets consist of a porous fabric, for example polyester fabric, lightly coated with vinyl. They are usually are used for sun and wind protection, but they do not protect from rain. Knitted nets are produced with high density polyethylene and polypropylene or acrylic fibres. The third type of membrane, are fabrics that are widely used. In fabric structures, the materials used are meshes, nettings and films. The mesh consists of a porous fabric with open spaces between the yarns and is produced with several types of fibre. One of the meshes comprises polyester weaves lightly coated with vinyl which have mechanical resistance, present lower water absorption, easy dying and are an inexpensive material. Polyamide fibres are frequently used for indus-
trial applications due to their properties, such as, high strength and chemical resistance, although they have high water absorption and are more costly than polyester. Netting can use polyester, polyamide or polypropylene filaments or spun yarns which are knitted or knotted in order to produce the fabric. Each of these fibres presents advantages and disadvantages. For instance, polyester holds better than polyamide, it is not so easy to coat and is more expensive. Polyamide is easier to coat but has a higher water absorption rate than polyester and does not hold the dye as easily. Regarding films, they are transparent polymers extruded in sheet form without any supporting substrate. Some examples are vinyl, polyester or polyethylene. Although they are not as expensive as textile structures, they have several drawbacks, namely less strength and durability and more elasticity than textile structures, making them not a very suitable option for several applications [59, 61].

For architectural applications, fabrics need to have special properties and special forms that cannot be chosen randomly. Thus, the production of textile architectural membranes is divided into two phases: the conceptual design and analysis of the structures and the use of advanced geometry in the production of the built form. These fabrics present a totally different behaviour than the traditional ones, being characterized by non-linear, anisotropic and non-elastic behaviour. The fabrics are also resistant to abrasion and mechanical degradation, impermeable to water, air and wind and resistant to the sun and acid rain degradation when exposed during long periods of time. The fibres are resistant to deformations and extensions under loads, characteristically obtained by the shape given to the fabric. The fabrics should have sufficient curvature to enable the tensile forces applied. They need also to be lightweight, flame resistant, present a long life service and be translucent. The low mass and the translucency are two properties which contribute to thermal comfort inside the building system [59, 63, 64].

To obtain a good textile architectural structure that works according to the requirements, it is necessary to select the right material, the proper design, engineering, production and installation. Most textile structures use fabrics rather than meshes or films. In tensile membrane structures, the most important material is obviously the fabric.

Several fibres can be used in membranes, being their selection made according to the membrane end use. Some fibres present the appropriate properties having the potential to be applied; however, the high costs prevent their wide utilization. From all the synthetic fibres, the most used for membranes are polyester, glass fibre and aramid; other types of fibres are used in very specific applications, such as polyamide and cotton. Table 18.2 presents some of the properties of the mentioned fibres [60, 61, 62].
Polyamide is not preferred for this application due to its poor resistance against UV light and, when wet, polymide swells in length direction, reducing its interest to textile architecture. Polyamide textile structures could be more durable than polyester; however, a high cost and stretch are major drawbacks. For these reasons, polyamide textile structures should only be used for specific applications [62, 63].

Polyester fibre is considered as a standard product in this area, and together with glass fibre, is the most common fibre in textile architecture. The main reason for its wide use is directly related to its advantages, such as, good tensile strength, high durability, low cost, stretch, elasticity, and allows small corrections during installation. However, when exposed to sunlight, polyester mechanical properties decrease over time. Polyester fabrics are usually coated or laminated with PVC films resulting in a cheaper product (Figure 18.17).

The Austrian Company Sattler Ag is one of many companies manufacturing products for textile architectural applications using polyester yarns for base material. Their textile structures are used for pavilions for international trade fairs and exhibitions, halls and tents (e.g. for events and family parties), leisure centres, inflatable structures (tennis, pools and golf) and tailor-made structures. Some of the characteristics of Sattler’s material are presented in Table 18.3. According to Sattler Ag, the excellent

<table>
<thead>
<tr>
<th>Fibre</th>
<th>Tensile strength (N/mm²)</th>
<th>Tensile strain (%)</th>
<th>Elasticity (N/mm²)</th>
<th>Other properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyamide 6.6</td>
<td>Until 1000</td>
<td>15–20</td>
<td>5000–9000</td>
<td>• When exposed to light only average resistant to ageing; • Swelling when exposed to moisture; • Only little importance to textile architecture</td>
</tr>
<tr>
<td>Polyester</td>
<td>1000–1300</td>
<td>10–18</td>
<td>10000–15000</td>
<td>• Widely spread • Standard product • When exposed to moisture, reduction of breaking strength</td>
</tr>
<tr>
<td>Glass fibre</td>
<td>Until 3500</td>
<td>2.0–3.5</td>
<td>70000–90000</td>
<td>• Brittle fibres • Standard product • Special fibre for high-tech products • Costly</td>
</tr>
<tr>
<td>Aramid</td>
<td>Until 2700</td>
<td>2.0–4.0</td>
<td>130000–150000</td>
<td></td>
</tr>
</tbody>
</table>

Table 18.2 Properties of fibres used in textile architectural membranes
material used presents greater rigidity than steel, low maintenance, tear, ageing and weather resistance combined with lightness and elegance [66].

As far as the laminating is concerned, vinyl is applied to the woven or knitted polyester meshes, usually known as scrims or substrates. Normally, the textile structure base used is characterized by a high count and high tensile. The laminate provides extra strength. One of the production methods of polyester textile structures allows the coating during its production. All process consists in the production of the fabric that is placed under tension before and during the coating process. The result is a fabric with good dimensional stability once the yarns, in warp and weft directions, present similar characteristics. Depending on the fabric characteristics, it is possible to have different applications. For instance, usually lighter fabrics are used for acoustic and thermal insulation liners suspended beneath a structure’s envelope. However, if the application is for exterior, demanding a long service lifetime, heavier fabrics are most suitable. In this case, above the coating is applied a layer named topcoating, for instance, polyvinyl fluoride (PVF) or polyvinylidene (PVDF). Both types of topcoatings provide a protective finish to withstand environmental degradation. To prevent the hardening and the degradation by sunlight and fungus attacks and to provide fire retardancy, several additives are added to the material. An acrylic and PVDF finish are two types of additives. The first provides a better service life and the second an improvement to the soil behaviour [61, 63, 64, 65].

PVC coated polyester membranes present the following characteristics [67, 68, 69]:

- one of the least expensive;
- design life between 15 and 20 years;
- robust and durable;
- easier to ship and erection can be mountable;
Table 18.3 Sattler products characteristics. Examples [66]

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Unit</th>
<th>Type I</th>
<th>Type VI</th>
<th>745</th>
<th>787 114</th>
<th>787 181</th>
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<tbody>
<tr>
<td>Product</td>
<td></td>
<td>648</td>
<td>717</td>
<td>Uni</td>
<td>Antiwicking</td>
<td>Opaque</td>
</tr>
<tr>
<td>Coating</td>
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<td>Acrylic/PVDF</td>
<td>Acrylic/PVDF</td>
<td>Acrylic/PVDF</td>
<td>Acrylic/PVDF</td>
<td>Acrylic/PVDF</td>
</tr>
<tr>
<td>Weave</td>
<td>L 1:1</td>
<td>L 3:3</td>
<td>L 1:1</td>
<td>L 1:1</td>
<td>L 1:1</td>
<td>L 1:1</td>
</tr>
<tr>
<td>Yarn</td>
<td>dTex</td>
<td>1100</td>
<td>1670</td>
<td>1100</td>
<td>1100</td>
<td>1100</td>
</tr>
<tr>
<td>Ends/picks</td>
<td>9/9</td>
<td>14/14</td>
<td>7/7</td>
<td>8/8</td>
<td>8/8</td>
<td>8/8</td>
</tr>
<tr>
<td>Total weight</td>
<td>g/m²</td>
<td>700</td>
<td>1350</td>
<td>590</td>
<td>670</td>
<td>750</td>
</tr>
<tr>
<td>Tensile strength</td>
<td>N/5 cm</td>
<td>3300/3000</td>
<td>7500/7000</td>
<td>2100/2300</td>
<td>3000/2800</td>
<td>3000/2800</td>
</tr>
<tr>
<td>Flame retardency</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>
translucency can reach up to 20% depending on the thickness of the material;
• wide range of possible colours;
• flame retardant;
• easily cleaned;
• good tensile strength and dimensional stability.

This kind of coating allows the use of different chemical additives that guarantees the leakage proofness, light transmission, UV resistance and abrasion resistance, among others. With these features it is possible to have properties like resistance to fungus and bacteria, temperature resistance and antistatic properties [65].

Vinyl-coated polyester membranes are usually used for producing flexible structures, like awnings, canopies, walkways, tent halls, smaller air-supported structures and light member-framed structures. This membrane presents a polyester scrim, a bonding or adhesive agent and an exterior yarns which confer to the membranes excellent dimensional stability allowing bending without losing any tensile properties. The function of the polyester scrim is to support the coating and provide the tensile and tear strength, elongation and the dimensional stability of the final fabric. Polyester fibres are bonded to the coating by an adhesive agent. The adhesive agent, besides the bonding function, prevents the wicking of moisture into the fibres. Properties such as, water resistance, mildew resistance, flame retardancy, different levels/degrees of translucency (from high levels of light transmission to completely opaque) and a wide range of colours, characterize the vinyl-coated polyester.

Polyester mesh fabric produced by Jiangsu Jiuding New Material Company is a reinforcement, which is an industrial fabric made by a warp knitted using polyester yarn. This product is used as base fabrics for vinyl laminated fabrics and coated fabrics. Applications are usually tent and structural material and awning and canopy material among others. Polyester mesh fabrics present properties such as, high tensile strength and weight, extensibility, conformability, long-term dimensional stability and can be widely used, which make them excellent products for the applications previously mentioned. Some of the products available are listed in Table 18.4 [70].

Awnings, tents and low tension frame structures are applications where vinyl-laminated polyester can be used. Laminated fabrics consist normally in two or more layers of fabric or films bonded together with the help of an adhesive agent by a heat or pressure process. A good bonding is required once it is necessary to have the proper seam strength and to prevent delamination. The use of an open-weave scrim, for instance a mesh, makes
these fabrics more economical. But if colour, UV resistance, abrasion resistance are required, then the product final price will be high. Vinyl-laminated polyester is a good option for backlighting [61].

A membrane is a unique structure, lightweight and flexible, that presents higher strength/weight ratio when compared with concrete or steel. It is environmentally friendly and in most cases, the fabrics are recycled, and can be designed for almost any condition. All these features are advantages for the textile structures. The low rigidity of these structures could be a problem, thumb spans greater than 15 m should be avoided, loss of tension can result in failure of the structure, there are thermal limits, and in an open system the water control is difficult and thus needs special attention.

18.5.2 Applications

With fabrics made of high tensile coated polyester, a whole new application comes up in the field of building. Due to its unique properties like high flexibility, high tensile strength, the lightweight and the high light transmission ratio of the material it is possible to apply this structure to, ‘roof large without trusses, to design spaces flooded with light, structurally realise completely new form, to generate attention by design and colour, to establish temporary structures and to set them up again at another location, to build in a resources-saving manner and to develop completely new building areas’ [65].

The applications of textile architectural membranes for building include a wide range of areas, such as, sports and leisure, arts and culture, travel and transportation, business and commerce, and agriculture and environmental industries. Some examples of those applications are [65, 69]:

- Public event building: using temporary or permanent roofing and also enclosed building forms for exhibitions, shows, among others (Figure 18.18).
Tourism and catering: usually used for sun, wind and rain protection providing a comfortable environment; also their lightness and expanse are attractive characteristics (Figure 18.19).

Parks and landscape spaces: protect the visitors from the weather conditions and are generally suited to the natural environment (Figure 18.20).

Open-air theatres: during this type of event protection against the weather is required (Figure 18.21).

Sports buildings: the main function is the weather protection; one of the most important features is the high light transparency of the membranes (Figure 18.22).

Entrance and walkway areas: besides the technical function (provides protections against weather conditions) also works as a decorative element (Figure 18.23).
18.20 Park architectural membranes [65].

18.21 Open-air theatre architectural membranes [65].

18.22 Sports building architectural membranes [65].
Shopping centres and exhibition spaces: exceptional membrane architecture supports marketing concepts which make shopping experience-oriented (Figure 18.24).

Railway and bus stations: the main function, in this case, the protection although aesthetic function is also an aspect which is usually considered (Figure 18.25).

In addition, there are classical applications and others that can be done with different functions, such as:

- for heat- or noise-insulating purposes, a multilayered membrane is used;
- for light protection, i.e. opaque membranes for specific requirements like cinema halls and some greenhouses where the light should not enter in the building;
for textile secondary structures for shading sails;
- for interior architecture as a decorative element;
- for applications that only requires demountable, temporary structures for short-term.

Ceno Tec is an Austrian company that offers other innovative applications, such as: flexible bulk containers like big bags (Figure 18.26), protective cloth/hoods and oil retentions/cascade separators (Figure 18.27).

Textile architectural membranes are widely used and for many and different reasons. Their unique properties allow them to have the flexibility to improve and lead them to innovative technologies. Further developments in membranes will be related to the materials development and its application. New applications and materials development ensures the use of membranes in the architecture of the future [58].
18.6 Ocean engineering applications

Besides the previous applications, namely concrete reinforcement, geotechnical, environmental and textile architectural, polyester and polyamide fibres are also used in a specific area of civil engineering field, ocean engineering.

18.6.1 Civil engineering in the ocean

Civil engineering embraces a wide variety of technical areas in both theoretical and experimental research. It deals with different areas such as, planning, design, construction, repair, maintenance and management of physical infrastructure networks [71, 72].

One of the civil engineering branches is ocean engineering, a hybrid technical area, which is a relatively young and remarkably exciting field of engineering. It is the application of basic engineering to the analysis, design, construction and management of systems that operate in the ocean environment. Some of the areas which are covered and included in this discipline are: Offshore Engineering; Naval Architecture; Marine Structural Mechanics; Safety and Reliability; Materials; Cable; Mooring, Buoy Technology; Underwater Technology; etc. [73].

According to the Society of Naval Architects and Marine Engineers, ‘ocean engineers study the ocean environment to determine its effects on ships and other marine vehicles and structures’ [74]. Ocean engineers may design and operate stationary ocean platforms, or manned or remotely operated sub-surface vehicles used for deep sea exploration. They have petroleum engineering tasks such as underwater oil or gas exploration, or they might design structures such as offshore drilling platforms, harbour facilities, and underwater machines [75].
One of the most important aspects of ocean engineering concerns how environmental conditions such as, wind and wave climate, currents, seawater, hydrostatic pressure and earthquakes, affect the different types of civil structures including the equipment used, and how the structures, their foundations, and equipment respond to these factors [72].

18.6.2 Fibre ropes for construction engineering in the ocean

Rope technology has begun with organic fibre material which was spun and twisted into a larger more useful form without any binding element [76, 77]. Rope modernization started with the development of polyamide fibres (nylon) in 1938, followed later with the introduction of other synthetic fibres, such as polyester [76]. In recent decades advances have been made in this field which greatly influenced civil engineering in the oceans during this century. Fibre technology innovations led to the development of new fibres with different properties and performances for a wide range of applications. One of those applications includes the construction and reinforcement of ropes and cables, widely used in this field of civil engineering [78]. Thus, for conventional rope-making, the most common fibres are polyester, polyamide and polypropylene. Table 18.5 presents the general properties of fibres used in the production of ropes as well steel wire [79].

Conventional steel wire mooring systems present several problems as water depth increases, namely, lower restoring efficiency, high proportion

<table>
<thead>
<tr>
<th>Fibre</th>
<th>Specific modulus (N/tex)</th>
<th>Specific strength (N/tex)</th>
<th>Other characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyamide</td>
<td>4</td>
<td>0.84</td>
<td>10–15% wet strength loss</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Poor wet internal abrasion resistance</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Moderate creep</td>
</tr>
<tr>
<td>Polyester</td>
<td>10</td>
<td>0.84</td>
<td>Good wet internal abrasion resistance</td>
</tr>
<tr>
<td>Polypropylene</td>
<td>8</td>
<td>0.73</td>
<td>Lighter than water</td>
</tr>
<tr>
<td>monofilament</td>
<td></td>
<td></td>
<td>Moderate creep</td>
</tr>
<tr>
<td>Polypropylene</td>
<td>3.5</td>
<td>0.66</td>
<td>Lighter than water</td>
</tr>
<tr>
<td>multifilament</td>
<td></td>
<td></td>
<td>Moderate creep</td>
</tr>
<tr>
<td>Steel wire</td>
<td>26</td>
<td>0.18</td>
<td>Typical properties</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Corrodes</td>
</tr>
</tbody>
</table>
of tether strength is consumed by the vertical components of line tension, reduced pay-load of the vessel, large mooring radius and sea-floor foot print, costly to maintain, mechanically complex, relatively inflexible and it is too heavy. All these disadvantages prompted engineers to find a substitute material. Thus, fibre ropes seem to be an interesting alternative once they can solve some of those problems [80, 81]. Synthetic fibre mooring ropes are lightweight (20% compared with wire rope) and more flexible, thus being much easier to handle. Synthetic fibre ropes do not corrode with the salt-water environment; have high tensile strength and adequate axial stiffness similar to wire ropes. However, they are much more susceptible to damage than steel wire. The ropes’ elasticity allows them to absorb the energy of the movements under adverse environmental conditions (i.e. excessive wave and winds), and they possess excellent resistance to abrasion, wear, creep and thermal degradation [76, 82, 83]. Generally synthetic fibres can be used in mooring systems, in anchoring and in logging applications [82, 84, 85].

There are three types of rope construction which can be used: plaited, twisted and braided. The rope design and construction are important issues since rope performance depends on them. Each construction provides different properties [86, 87]:

- **plaited**: resists rotation; does not kink or twist; has good strength and elongation; good weight;
- **twisted**: offers good strength; easy handling; tendency to ‘unravel’ when placed under load; may cause failure;
- **braided**: higher in strength and durability; lower in elongation; very pliable and easy to handle; more difficult to splice; higher cost; single and solid – braided types are more reliable than double-braided (Figure 18.28).

![Braided rope type (Cordoaria Oliveira Sá, Portugal).](image)
Polyamide fibre has been used in marine mooring since the 1950s approximately. This fibre is characterized by low modulus presenting the lower value from all quoted fibres, and is the strongest one when dry. However, when wetted, it loses 10% of its strength although that value can go up to 20%. In these conditions, polyamide also suffers strength loss due to creep and internal abrasion during cyclic tensile loading. All these disadvantages, along with the short service life, make polyamide ropes, in a general way, unsuitable for permanent deep water moorings.

Notwithstanding that fact, nowadays there are excellent ropes made of polyamide yarns. For instance, Enka®Nylon is a high-tenacity polyamide yarn used for mooring ropes with the ability of high shock absorbance and long time reliability. The wide range of high-tenacity polyamide yarns with Enka®Nylon is used in several fields of technical applications where safety and reliability are key requirements. One of those areas is ocean engineering. Enka®Nylon 140HRT and Enka®Nylon 142HRT are two types of polyamide yarns usually used for the production of mooring ropes. Some of their characteristics are described in Table 18.6 and Table 18.7 [88]:

### Table 18.6 Enka®Nylon 140HRT characteristics [88]

<table>
<thead>
<tr>
<th>Linear density (nominal dTex)</th>
<th>Number of filaments (nominal)</th>
<th>Linear density (dTex)</th>
<th>Breaking strength (N)</th>
<th>Breaking tenacity (mN/Tex)</th>
<th>Extension at break (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>700</td>
<td>108</td>
<td>716</td>
<td>60.7</td>
<td>848</td>
<td>19.5</td>
</tr>
<tr>
<td>940</td>
<td>140</td>
<td>942</td>
<td>79.5</td>
<td>844</td>
<td>17.9</td>
</tr>
<tr>
<td>1400</td>
<td>210</td>
<td>1405</td>
<td>118.6</td>
<td>844</td>
<td>18.9</td>
</tr>
<tr>
<td>1880</td>
<td>280</td>
<td>1895</td>
<td>159.2</td>
<td>840</td>
<td>19.5</td>
</tr>
<tr>
<td>2100</td>
<td>280</td>
<td>2105</td>
<td>175.1</td>
<td>832</td>
<td>19.8</td>
</tr>
</tbody>
</table>

### Table 18.7 Enka®Nylon 142HRT characteristics [88]

<table>
<thead>
<tr>
<th>Linear density (nominal dTex)</th>
<th>Number of filaments (nominal)</th>
<th>Linear density (dTex)</th>
<th>Breaking strength (N)</th>
<th>Breaking tenacity (mN/Tex)</th>
<th>Extension at break (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>940</td>
<td>140</td>
<td>938</td>
<td>90.0</td>
<td>960</td>
<td>16.9</td>
</tr>
<tr>
<td>1400</td>
<td>210</td>
<td>1407</td>
<td>133.4</td>
<td>950</td>
<td>18.4</td>
</tr>
</tbody>
</table>
Due to polyester fibres’ advantages, ropes made from these fibres are replacing polyamide ropes in many critical conventional marine applications. Technological innovation allowed the development and production of high quality polyester fibres enabling the production of strong polyester ropes with relatively high modulus. Very durable in cyclic tensile fatigue loading, stronger than polyamide in dry conditions, keeping strength and generally stronger than polyamide when wetted, are advantages which make polyester a good candidate for deep water mooring systems. On some occasions, polyester fibres are combined with polypropylene fibres because, although they present similar strength and stiffness properties to those of all polyester ropes, they have lower weight and cost, have great resistance to surface abrasion and heat build-up than all polypropylene ropes [79]. Nowadays, technology allows the production of high tech polyester yarns which can be used in ocean engineering in several operations. Marlow Ropes Ltd is a British company which produces several types of ropes for different areas, namely for marine applications like mooring and anchor lines. Three strand polyester is a rope made of high tenacity polyester yarns possessing special features. No strength loss when wet, good abrasion resistance, flexible and soft handle and easy splice are some of them. However, they present other properties which make them suitable for the applications mentioned, such as, UV resistance (50% strength retention after three years) and good resistance to acids and alkalis. Tables 18.8 and 18.9 show some of the characteristics of the three strand polyester rope.

The traditional practice of mooring with ropes, for thousands of years, remained unchanged till recently. A mooring system can be defined as a compliant, usually tensile structure, which restrains a vessel against the forces of the sea and the movements generated by natural forces such as wind, wave and currents. The mooring consists of a fixed mooring system.

### Table 18.8 Technical data of three strand polyester [89]

<table>
<thead>
<tr>
<th>Fibre</th>
<th>High tenacity polyester</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV resistance</td>
<td>50% strength retention after three years</td>
</tr>
<tr>
<td>Standard lengths</td>
<td>100–200 m reels</td>
</tr>
<tr>
<td>Recommended splice</td>
<td>Easily spliced using three strand method</td>
</tr>
<tr>
<td>Chemical resistance</td>
<td>Good resistance to acids and alkalis</td>
</tr>
<tr>
<td>Colours</td>
<td>White, black and navy blue</td>
</tr>
<tr>
<td>Sheave diameters</td>
<td>10:1 or ten times rope diameter</td>
</tr>
<tr>
<td>Working loads</td>
<td>Maximum loads should not exceed 50% of the break load published</td>
</tr>
<tr>
<td></td>
<td>Normally exceed 20–30% of the break load published</td>
</tr>
<tr>
<td>Relative density to H₂O</td>
<td>1.38 g/cm³</td>
</tr>
<tr>
<td>Critical temperatures</td>
<td>Melts at 260°C</td>
</tr>
</tbody>
</table>
that can be platforms, cells, spuds and bollards, a pile system, and a mooring line system which includes natural or synthetic fibre rope, wire rope and mooring chains (Figure 18.29). As can be seen in Figure 18.30, floating mooring systems can be installed in three different ways: tension leg (A), taut leg (B) and catenary (C) [90, 91].

The best structural concept for synthetic fibre moorings is the taut leg system. This allows polyester rope to be deployed from the platform at

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approximately 45° from vertical and placed under tension. Many are the advantages in using the mooring design taut leg for polyester fibres, namely [76, 80, 82, 85]:

- low creep and good creep rupture resistance;
- good fatigue performance;
- low sensitivity to alternate tension compression;
- low hystereses;
- elasticity of synthetic line provides the restoring force, i.e., there is a favourable force vector to restore the platform to its neutral position;
- reduction of mooring line weight hence better vessel payload;
- more efficient system allowing a smaller footprint in the ocean;
- lower cost;
- due to its lower tensile stiffness an extreme reduction is visible/verified in line dynamic tension;
- very low rate of hydrolysis;
- corrosion resistance;
- easy handling and subsequent simpler and better installation techniques;
- in most cases lower vessel offset.

Fibre ropes have been used for more than 30 years in offshore and ocean industry, but polyester ropes in permanent moorings are relatively new. However, they have been extensively studied and tested in field and laboratories and, nowadays, polyester ropes seem to be well established in deepwater moorings [82]. There are several possible failures in ropes, categorized as: environmental, surface, wear tensile and structural failures (like tensile overload or torque effects) and fatigue. One of the huge prob-
lems with steel is the corrosion due to the sea-water, but the same occurs in polyester ropes because of hydrolysis, although less significant. Environmental effects like UV and microbiological attack are not a concern since polyester has an excellent resistance to both [76]. Rope installation needs to be done carefully because most of the damage occurs by mishandling during this stage, in which ropes can be cut, crushed or suffer abrasions against objects [76, 82]. Tensile overload should not occur if the rope is properly designed, although severe twisting or flexion should be avoided. In polyester fibres, the fatigue due to transverse crack growth, under cyclic loading, is not a failure mechanism as it is for steel. According to Hearl et al., there are five types of fatigue effects which need to be considered for ropes: tensile fatigue, hysteresis heating, creep rupture, internal abrasion and axial compression. The tensile fatigue, which should be up to 20% minimum load at 70% maximum, is almost nonexistent in polyester fibres. The same happens regarding creep rupture that occurs under static and dynamic cyclic load; it would be necessary to produce it over many thousands of years. Hysteresis heating is not significant for strain amplitudes less than ±0.5%, however this could be a problem with larger amplitudes which may occur in shallow water moorings. For polyester fibres, internal abrasion only takes effect after millions of load cycles due to reduced relative fibre movement, but it is a serious problem for highly twisted polyamide ropes in wet conditions. Due to its low modulus, axial compression fatigue, according to Flory [79], is not likely to be a problem for polyester fibres, being necessary for a large number of cycles to occur. Also they can support thousands of very low trough load cycles or a complete relaxation without losing significant loss.

Based on the studies already undertaken in yarn fatigue mechanisms, one can conclude that polyester ropes designed according to current standards and planned lifetimes, give no reason to expect failures, which make them a very good option for mooring systems. Due to absence of corrosion and of metal fatigue, polyester rope performance should be better than steel ropes. Besides, if well designed, axial compression fatigue does not occur and internal abrasion appears to be the cause of the loss of strength after millions of cycles. However, the fatigue performance of polyester ropes depends on different factors like, fibre quality, rope construction, termination design and on cycle load history [76, 78, 82, 86, 92].

The key to a successful anchoring system is the structure which attaches the vessel to the anchor. Two options are available to anchor, wire ropes and natural or synthetic ropes. The first option presents two distinct advantages, brute strength and its weight. However, the absence of elongation, the difficulty in handling it due to its weight and the transmission of heavy shock-loads to the vessel and the anchor make them not very good for this operation. Regarding natural fibre ropes, there are several drawbacks
namely: low elongation, high weight and high bulk, no resistance to the effects of rot, sun damage and shipboard vermin. They also transmit heavy-shock loads to vessel and anchor [85].

When synthetic ropes were introduced in the ocean industry, polyamide was one of the fibres which were used for anchoring applications (Figure 18.31). By that time, its uses gave a new dimension to the sector, becoming extremely important because polyamide performance was better than wire and natural fibre ropes. Polyamide ropes are strong, resistant to sun and rot damage and, above all, they have elasticity allowing stretching without breaking and absorbing a part of the load applied on the anchoring system, for instance winding and wave action, maintaining the anchor in the right position. One of the most common construction methods of ropes using polyamide is called three-strand nylon anchorline, usually used to anchor a vessel in an adequate way being characterized by a reduced diameter rope.

Comparing polyester fibre with polyamide, the latter absorbs a considerable amount of water leading to significant changes in their properties. After wetting and once the polyamide fibres become dry, their handling becomes very difficult with resistance to flaking or coiling. Moreover, polyamide presents an aliphatic linear structure while polyester has an aromatic ring which gives a higher stiffness to the fibre [79, 85]. After many cycles of dynamic loads, a decrease in physical strength is observed in polyamide fibres which can lead to failure. Due to the twisting given during production to typical three-strand polyamide rope, they have tendency to kink or hackle especially when used with mechanical winches.
Yale Cordage developed a new manufacturing rope with polyamide fibre, eight-strand rope, with the same strength and abrasion resistance to the polyester version, which can solve many of the above mentioned problems [84] (Figure 18.32). This unique construction process allows that less storage is required, lower pull-out force due to superior energy absorption, superior performance in power windlasses and non-hackling and torque-balanced performance.

According to Lee et al. [81], considering all possible materials and production techniques that can be used for rope production there is not enough experience to determine which is the best for mooring systems and other applications. Since synthetic fibre rope properties will influence the system performances, it is necessary to consider the design and the system analysis, rope design, rope specification and testing, rope production and quality assurance, rope handling and installation and in-service inspections and maintenance of the ropes applied in the ocean industry [81, 85].

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