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Membranes for Industrial Wastewater Recovery and Re-use
Membranes for Industrial Wastewater Recovery and Re-use
To Claire, Oliver and Samuel
Membranes for Industrial Wastewater Recovery and Re-use

Edited by:
Simon Judd and Bruce Jefferson
Contents

Preface ix
Contributors xiii

Chapter 1 Introduction
1.1 1.1 Water reuse motivations and barriers 2
1.2 Industrial water 4
1.3 Membrane technology 7

Chapter 2 Membrane technology
2.1 The membrane 14
  2.1.1 Membrane and membrane process definition 14
  2.1.2 Membrane structure 14
  2.1.3 Membrane materials and their manufacture 17
  2.1.4 Membrane configurations 23
2.2 The process fundamentals 32
  2.2.1 Process performance definitions 32
  2.2.2 The driving force 34
  2.2.3 Factors opposing the driving force 34
  2.2.4 Critical flux 39
2.3 The theory 40
  2.3.1 Membrane mass transfer control 41
  2.3.2 Fouling/cake layer mass transfer control 43
2.4 Process design and operation 52
  2.4.1 Staging 52
  2.4.2 Specific energy demand 55
  2.4.3 Fouling and pretreatment 59
  2.4.4 Backwashing and cleaning 66

References 70
Chapter 3 Industrial waters

3.1 The power industry: water requirements for power generation and cooling water
   3.1.1 Water demand by the power industry
   3.1.2 Overview of cooling water systems
   3.1.3 Overview of boiler feedwater systems
   3.1.4 Sources of water used for
   3.1.5 Water quality comparisons
   3.1.6 Optimisation of water use in recirculating cooling systems
   3.1.7 Cooling tower water quality issues
   3.1.8 Governing legislation and guidelines
   3.1.9 Volumes and quality of aqueous process waste streams
   3.1.10 Current reuse practices and opportunities

References

3.2 The pulp and paper industry
   3.2.1 Introduction
   3.2.2 Pulping and paper manufacturing processes
   3.2.3 Effluent volumes and quality
   3.2.4 Current water and effluent purification systems and governing legislation
   3.2.5 Membranes in the pulp and paper industry
   3.2.6 Conclusions

Acknowledgements

References

3.3 The textile industry
   3.3.1 Categories of textile processing operations
   3.3.2 Effluents from textile processing unit operations
   3.3.3 Process water quality requirements
   3.3.4 Legislation
   3.3.5 Conventional treatment
   3.3.6 Demand management
   3.3.7 Reuse practice and opportunities

References

3.4 The beverage industry
   3.4.1 Point of use recycling opportunities
   3.4.2 End of pipe recovery opportunities

References

3.5 Pure waters in the pharmaceutical industry
   3.5.1 Background
   3.5.2 Water quality standards
   3.5.3 Volumes and quality of aqueous process and waste streams
   3.5.4 Reuse opportunities
Chapter 4  System design aids

4.1  Computer-aided design for reverse osmosis plant  172
   4.1.1  Introduction  172
   4.1.2  Key elements of the reverse osmosis process  172
   4.1.3  RO design software  176
   4.1.4  Cost calculation  182
   4.1.5  Overview  183
   References  186

4.2  Water pinch analysis  186
   4.2.1  Introduction  186
   4.2.2  Water pinch: the history  187
   4.2.3  Methodology  187
   4.2.4  Computed solutions  191
   4.2.5  Software tools currently available  197
   4.2.6  Case study: water pinch and implementation of regeneration techniques  201
   4.2.7  Conclusion  209
   References  209

4.3  Design examples  213
   4.3.1  Problem in reverse osmosis: film theory and energy demand  213
   4.3.2  Problem in reverse osmosis: array design  216
   4.3.3  Problem in reverse osmosis: CAD array design  217
   4.3.4  Problem in electrodialysis: energy demand  220
   4.3.5  Problem in submerged membrane filtration design  223

Chapter 5  Case studies

5.1  Flag Fen high-purity water production plant (UK)  228
   5.1.1  Background  228
   5.1.2  Description of plant  228
   5.1.3  Performance  230

5.2  Eraring Power Station: purification of secondary sewage for boiler feedwater (Australia)  232
   5.2.1  Background  232
   5.2.2  Description of plant  233
   5.2.3  Performance  235

5.3  Doswell combined cycle power plant: zero liquid discharge (USA)  237
   5.3.1  Background  237
   5.3.2  Description of system  238
   5.3.3  Performance  240

5.4  VHP Ugchelen: paper mill water recycling (Netherlands)  241
   5.4.1  Background  241
   5.4.2  Description of system  242
In setting out to produce a book on any given subject, the primary questions to be answered are: "What should we write about?" and "Who's going to buy it?". These questions are obviously interlinked.

The subject area of this book is well defined in the title: membrane technology and its application to industrial water recycling. There can be little doubt of the importance of recycling of water contaminated by industrial activity, and it is hoped that the introduction to this book (Chapter 1) sufficiently emphasises this point. Stresses imposed on freshwater supplies continue to grow and environmental legislation relating to discharges becomes ever more rigorous. Given that technologies for water purification are tending to become more efficient and generally cost-effective, it is inevitable that recovery and reuse of effluent will be more widespread in the future. It is also the case that membrane processes play a pivotal role in many reclamation and reuse schemes in industry. Their application across all industrial sectors has increased exponentially over the last twenty years and there is little sign of this growth abating.

Since the subject is very extensive the amount of detail is largely determined by the target readership, which relates to the second key question. This is rather vexing matter, not least because potential readers are well served by a number excellent texts on membrane processes and technologies (Table 1) as well as a growing number on water reclamation and reuse (Table 2). Neither of these tables is comprehensive, and it can always be argued that there is really nothing new to write about.

So, who might buy this book? What function is this book meant to serve? Perhaps it is as well to outline what the book is not. The book is not meant to be a comprehensive treatise on membrane processes and technologies, although an outline of these areas is provided (Chapter 2) with reference to membrane materials, modules, and processes along with governing equations for use in system design. For more comprehensive information than that provided the interested reader may seek out such tomes as those produced by Ho and Sirkar, Rautenbach and Albrecht, and Scott (Table 1). Moreover, many of the industrial process applications in which membranes find employment, such as the biotechnology and food industries, are only briefly discussed with reference to a
few specific industrial sectors (Chapter 3). Again, industrial process applications are reasonably well documented in Cheryan (for filtration processes), Amjad (reverse osmosis) and otherwise the general texts previously mentioned.

It is also the case that the book does not provide much detail of domestic or municipal schemes, where water is either recycled within buildings or reclaimed from municipal wastewater treatment plant and reused for extensive schemes such as irrigation and aquifer recharge. There is also no discussion of sustainability as a concept, or any of the related topics pertaining to environmental impact and more holistic topics. Again, the interested reader is
already very well served in such areas, and in particular extensive systems, by the seminal work of Asano and the recent eclectic publication by Lens et al. (Table 2). Finally, although the principles of water pinch analysis are described in Section 4.2, this is only to enable an understanding of how it can be used. Once again, there are other texts available that comprehensively describe the methodology and its derivation (Mann and Lui).

It is hoped by the editors, however, that anyone interested in industrial wastewater recycling, including problem holders and membrane suppliers as well as students and academics studying in this area, will find this book useful. In particular, the book is meant to provide a practical aid to those readers with an interest in actually selecting, installing and/or designing systems for recovering and reusing industrial effluent. To this end, there is one chapter dedicated to design (Chapter 4), and in particular computer-aided design, based on commercial packages, of reverse osmosis arrays (Section 4.1) and water networks (Section 4.2). There are also eleven individual case studies, covering a range of applications, which are discussed with reference to design and operational facets of the water recycling treatment schemes (Chapter 5). Whilst not all industrial sectors where water recovery and reuse takes place are covered in this book (it is extremely difficult to extract all information pertaining to industrial water quality, let alone water reuse) as much detailed as has been made available is reported in the book. Lastly, the book is very much biased towards engineering practice, rather than scientific study. Data pertaining to membrane process performance in Chapters 3 and, more especially, Chapter 5, are based almost entirely on existing pilot or full-scale systems, rather than bench-scale studies or theoretical predictions.

As with any piece of work the editors would welcome any comments from readers, critical or otherwise, and our contact details are included in the following section.
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The editors

Simon Judd
Dr Judd is a Reader in Water Sciences at Cranfield University with over 15 years' experience of industrial and academic R&D, having spent three years in nuclear waste management and two years in forensic science. In his current post he has conducted research into a wide range of chemical and, principally, membrane processes as applied to water and wastewater treatment. Topics have included municipal and domestic wastewater treatment and reuse using membrane bioreactors, nanofiltration of dye waste for water recovery and scale mitigation in reverse osmosis for flux enhancement. He has contributed to and co-authored a number of books on both water recovery and reuse and membrane processes, and has published extensively in peer-reviewed journals. Dr Judd also manages swimming pool water treatment research activities, and chairs the Membrane Academia Industry Network (MAIN), a UK research council-sponsored initiative to encourage dissemination of information between the industrial and academic communities in the UK.

Contact: s.j.judd@cranfield.ac.uk

Bruce Jefferson
Dr Jefferson is a Senior Research Fellow at the School of Water Sciences at Cranfield University following graduation from the Department of Chemical Engineering, Loughborough University. In his current post he has conducted research into general physical and, more specifically, membrane processes for a wide range of applications within both urban and industrial environments. He has contributed to and co-authored a number of books on both water recovery and reuse and membrane processes, and has published extensively in peer-reviewed journals. Dr Jefferson has also presented papers at and chaired numerous sessions on the subject of water recycling, and he is a member of the IWA specialist group committee on advanced treatment/membrane processes.

Contact: b.jefferson@cranfield.ac.uk
The editors would like to convey their sincere thanks to the co-authors and other contributors to this book, details of whom are given below.

**The co-authors**

**Danielle Baetens**
Dr Baetens graduated in 1991 as a civil engineer in chemistry at the University of Ghent, obtaining a PhD at the faculty of Agricultural Engineering, also in Ghent, in 2000. The PhD concerned modelling of biological phosphorus removal in wastewater. Between 1991 and 1998 Dr Baetens worked at the University of Ghent as assistant at the faculty of civil engineering on the treatment of air pollution and wastewater treatment. Since 1999 she has worked at Vito (Flemish Institute for Technological Research), where she project is co-ordinator of the project group on process optimisation.
Contact: danielle.baetens@vito.be

**Etienne Brauns**
Etienne Brauns graduated as a Metallurgical Civil Engineer at KU Leuven in Belgium in 1976. He thereafter stayed at KU Leuven to contribute as a researcher to a program on shape memory alloys, joining the Belgian national nuclear research centre SCK-CEN in 1985. In 1990 all non-nuclear research at SCK-CEN was transferred to Vito, founded in that year by the Flemish government. As a researcher at Vito he originally continued R&D activities on technical and electro ceramics for sensor applications. He has subsequently focused on membrane research and currently works on pressure-driven and electrodialytic membrane processes, and water recycling.
Contact: etienne.brauns@vito.be
Mike Carney
Mike Carney qualified from Birmingham University with a First in chemical engineering and then undertook an MSc in water resources technology. He worked as a process design engineer for Paterson Candy and Schweppes where he was responsible for water and effluent treatment worldwide. As Technical Director of a water treatment contractor, he was responsible for the design of numerous food and beverage water and effluent treatment plants worldwide. Now working as an independent consultant at Vale Water Services Ltd, he specialises in the same food and beverage field. He is a Fellow of the Institution of Chemical Engineers.
Contact: vale.water@compuserve.com

John Hutcheson
Dr Hutcheson is currently General Manager of Pharmaceuticals at Christ Kennicott with extensive expertise in pure and ultrapure water treatment system design. On gaining a PhD in organic synthetic chemistry in 1980 he spent three years working in downstream processing at Beecham Pharmaceuticals followed by a short tenure at Millipore before becoming a founding member of Ionpure Technologies Ltd. He transferred to Kinetics in 2000, joining Christ Kennicott in 2002. He has authored a number of papers and presented at international conferences and courses on aspects of water purification.
Contact: john.hutcheson@christwt.co.uk

Audrey Levine
Audrey Levine is an Associate Professor of Civil and Environmental Engineering at the University of South Florida. She holds a BS degree in biology from Bates College (Lewiston, Maine), an MS degree in public health from Tulane University and a PhD in civil engineering from the University of California at Davis. Prior to joining the faculty at USF she was a faculty member at Utah State University (1994–1998), New Jersey Institute of Technology (1992–1994) and Iowa State University (1985–1992). She is actively involved in research and teaching related to wastewater reclamation and reuse, water distribution systems, potable water treatment, corrosion control and water quality issues. She is currently working on projects related to pathogen reduction through wastewater treatment, the fate of pharmaceuticals in reclaimed water systems and distillation of reclaimed water.
Contact: levine@eng.usf.edu

Jutta Nuortila-Jokinen
Professor Nuortila-Jokinen obtained her MSc in polymer chemistry at Helsinki University in 1988. Since 1987 she has worked in the Laboratory of MemPo, the Laboratory of Membrane Technology and Technical Polymer Chemistry at Lappeenranta University of Technology (LUT), as Research Assistant, Researcher and Associate Professor. She received her doctorate at LUT in 1997 before joining the Cleantech 2000 project (1997–2000) as Research Professor. She is currently working as Senior Fellow funded by the Academy of Finland.
(2000–2005). She has published around 60 scientific papers, and is a member of
the European Working Party on Membrane Science.
Contact: jnj@lut.fi

The contributors

In addition to the co-authors there are a number of contributors to this book who have provided specific information, in particular for the case studies detailed in Chapter 5.

<table>
<thead>
<tr>
<th>Name</th>
<th>Affiliation and address</th>
<th>Web address and contact</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alasdair Donn</td>
<td>PCI Membrane Systems, Laverstock Mill, Whitchurch, Hampshire, RG28 7NR, UK</td>
<td><a href="http://www.pcimem.com">www.pcimem.com</a> <a href="mailto:alasdair.donn@pcimem.com">alasdair.donn@pcimem.com</a></td>
</tr>
<tr>
<td>Robert P. Allison</td>
<td>Ionics, Incorporated, 65 Grove Street, Watertown, MA 02472, USA</td>
<td><a href="http://www.ionics.com">www.ionics.com</a> <a href="mailto:rallison@ionics.co.uk">rallison@ionics.co.uk</a></td>
</tr>
<tr>
<td>Devinder Chabba</td>
<td>Esmil Process Systems, Westfields, London Road, High Wycombe, HP11 1HA, UK</td>
<td><a href="http://www.esmil.co.uk">www.esmil.co.uk</a> <a href="mailto:sales@esmil.co.uk">sales@esmil.co.uk</a></td>
</tr>
<tr>
<td>Timo Sutela</td>
<td>Metso PaperChem Oy, PO Box 83, 21201 Raisio, Finland</td>
<td><a href="http://www.metso.com">www.metso.com</a> <a href="mailto:timo.sutela@metso.com">timo.sutela@metso.com</a></td>
</tr>
<tr>
<td>Hans Raemaekers</td>
<td>Triqua, Vadaring 7, PO box 132, NL-6700, AC Wageningen, The Netherlands</td>
<td><a href="http://www.triqua.nl">www.triqua.nl</a> <a href="mailto:hansramaekers@triqua.nl">hansramaekers@triqua.nl</a></td>
</tr>
<tr>
<td>John Murrer</td>
<td>Anglian Water Technology Group, Thorpe Wood House, Thorpe Wood, Peterborough, Cambridgeshire, PE3 6WT, UK</td>
<td><a href="http://www.anglianwater.com">www.anglianwater.com</a> <a href="mailto:jmurrer@anglianwater.co.uk">jmurrer@anglianwater.co.uk</a></td>
</tr>
<tr>
<td>Darren Morrison</td>
<td>Mitsubishi Corporation (UK) PLC, Mid City Place, 71 High Holborn, London, WC1V 6BA, UK</td>
<td><a href="http://www.mrc.co.jp/english/">www.mrc.co.jp/english/</a> <a href="mailto:darren.r.morrison@mitsubishicorp.com">darren.r.morrison@mitsubishicorp.com</a></td>
</tr>
<tr>
<td>David Holmes</td>
<td>Ultra tech, Wychbury House, 73c Stevens Road, Stourbridge, West Midlands, DY9 0XW, UK</td>
<td><a href="http://www.orelis.com/uk/home.orelis.asp">www.orelis.com/uk/home.orelis.asp</a> <a href="mailto:ultra.tech@btinternet.com">ultra.tech@btinternet.com</a></td>
</tr>
<tr>
<td>Hans Wouters</td>
<td>Paques Astraco, PO Box 52, 8560 AB Balk, The Netherlands</td>
<td><a href="http://www.paques.nl/">www.paques.nl/</a> <a href="mailto:h.wouters@paques.nl">h.wouters@paques.nl</a></td>
</tr>
</tbody>
</table>
Chapter 1

Introduction

Simon Judd
School of Water Sciences, Cranfield University
1.1 Water reuse motivations and barriers

The motivations for recycling of wastewater are manifold. Most often stated are those pertaining to increasing pressures on water resources. Reuse of wastewater conserves the supply of freshwater, and this presents clear advantages with respect to environmental protection. More pragmatically, wastewater reuse may result directly from legislation, which can constrain the discharge of polluted water by making this option onerous or else forbid such discharges altogether, or it may simply be favoured economically regardless of regulatory stipulations.

It is also the case that reuse itself is an emotive issue, and perhaps particularly so in the case of water. For domestic water recycling, that is recycling of water for non-contact domestic use such as toilet flushing or irrigation, public perception issues can outweigh the technical ones in terms of barriers to imposition. Key to this are the matters of ownership and identity. Studies have demonstrated that people are generally prepared to reuse water if it derives entirely from their own household, i.e. if they know where it has been. They are rather less prepared to use water if it is identified as deriving from some other source such as, for example, their neighbour's house (Jeffrey, 2002). Curiously, the complete loss of identity, such as arises either from large-scale community schemes, and indeed from conventional water supply via municipal works and intermediate environmental water bodies such as rivers, reservoirs and aquifers, is also perceived as being acceptable.

In reviewing the water reuse opportunities in industry, it is important to make the distinction between reclamation and recycling. Reclamation is the recovery and treatment of water to make it available for reuse; recycling is the recovery and reuse (whether or not subject to treatment) to and from a discrete operation. The development of water reclamation and reuse dates back centuries but modern day legislation probably dates back to 1956 in Japan, when the Industrial Water Law was introduced to restrict the use of groundwater by the rapidly growing Japanese industry, and Californian legislation leading to the adoption of the Federal Water Pollution Control Act of 1972 (Cologne, 1998). The drive for conservation of freshwater supplies has led to the development in some parts of the world, and in arid regions of the USA in particular, of large-scale community schemes in which water “recovered” from a municipal works is directly used for specific duties. In Japan, where sewerage services are limited and/or expensive, there has been a proliferation of in-building recycling schemes. For all such schemes, the key to the successful implementation of the reuse scheme is user acceptance and, ultimately, the assessment of risk.

Risk analysis, specifically to human health (Sakaji and Funamizu, 1998), plays a key function in municipal or in-building reuse schemes because of the implications of system failure for human health. Recycling of greywater (i.e. water used for washing) within a building, for example, is only likely to be acceptable if there is only a very minor risk to human health, predominantly from pathogenic microorganisms, arising from failure of the reuse system or some component of it. This imposes a limit both on the required rigour of the
repurification plant and, more directly, on the reuse options. Hence, reclaimed greywater for reuse in buildings is invariably employed exclusively for toilet flushing, where the risk to human health through exposure to the contaminated water is considered minimal. Large-scale municipal wastewater reclamation and reuse schemes are predominantly aquifer recharge, such as the well-established flagship Water Factory 21 plant in Orange County, California (Wesner, 1987) or turf grass irrigation schemes.

In the case of industrial water recycling the issue is notionally more straightforward. As long as recycling can reliably and cost-effectively provide water of a quality appropriate to its designated use, then there should be no barrier to its introduction and deposition of the existing freshwater supply. Of course, and as already mentioned, the cost benefit may be largely or wholly determined by statutory requirements: there are, for example, a number of power plants throughout the world that have a zero liquid discharge imposed upon their operation (Section 5.3). Under such extreme conditions, wastewater recovery and reuse is no longer an option but an absolute necessity — and inevitably incurs cost penalties due to solid waste generation and energy expenditure. In other cases, the decision to recycle is motivated entirely by economics; the total cost of purification to provide a water of a quality commensurate with or better than that of the freshwater supply is perceived as being less than the total cost of freshwater and waste discharge. Perception in this instance relates entirely to the acceptance of the technology, rather than to the use of water employed for duties considered unsavoury.

Given the apparently less restrictive constraints imposed on the recycling of industrial wastewater compared to that of domestic wastewater, it is perhaps surprising that the former has received far more attention, has attained a much higher profile and has achieved more widespread implementation than the latter. There exist a number of domestic and municipal wastewater recycling schemes, from single households through to large buildings, as well as municipal or industrial wastewater reclamation for irrigation. Similarly, a number of municipal wastewater reclamation schemes exist whereby the polished effluent is reused for cooling, which represents around two-thirds of all industrial water use, or other activities demanding low-grade process water such as for washing or transporting. Direct “closed-loop” industrial water recycling, whilst limited in technical and economic viability in many instances, is nonetheless attracting greater interest and being more widely applied. Indeed, it is already established practice for specific industrial processes where other resources are recovered in addition to the water, such as pulp solids in paper manufacturing (Sections 3.2.5 and 5.4–5.6) and paint pigments (Section 5.10) in electrophoretic painting.

Two factors mitigate against widespread “closed-loop” industrial effluent reuse. Firstly, most industrial processes involve a number of individual operations that give rise to wastewaters of a certain compositional range. These individual effluent streams are generally combined to give a wastewater whose resultant temporal variation in quality is immense, representing a significant challenge to any treatment process that is to provide water of a reliably high quality. Secondly, it is invariably the case that conventional sewage treatment
works have the capacity to treat these industrial waters simply by virtue of blending with domestic water, significantly dampening the effects of broad temporal variations in quality and the concomitant shock loads of specific problem contaminants that arise. In other words, existing municipal wastewater treatment works are often able to cope with industrial effluents at a cost that is considered reasonable. The loss of the industrial effluent biological and hydraulic load from a sewage treatment works designed to accept it may even cause operational problems. Moreover, a municipal water supplier is able to provide mains water of a quality consistent enough to be reliably employed in most industrial processes. A reduction in mains water consumption and effluent discharge has a negative impact on the revenue of the water supplier and sewer operator companies, who are often one and the same company. Given these implications, it is prudent to consider industrial water usage and discharge as a whole.

1.2 Industrial water

Industry accounts for about a quarter of all water consumption, and there is hardly any industry that does not use large volumes of water (Table 1.1). Although some industries abstract water from rivers and boreholes, much of the water used by industry is taken from public water supplies, and has therefore been treated to potable quality standards. This means that it is often of better quality with respect to microbial levels but nonetheless requires further purification to reduce the mineral and organic materials content according to the specific duty to which it is to be put (Tables 1.2 and 1.3). Although water consumption has actually decreased over the last 15 years in some regions of the world (Table 1.4), the price of supply and discharge has risen substantially over the same period. According to figures for the UK, the average cost of water supply and sewerage services to unmetered customers has more than trebled in the last

<table>
<thead>
<tr>
<th>Industry</th>
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<tr>
<td>Paper</td>
<td>29 m³/t paper produced</td>
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<tr>
<td>Newspaper</td>
<td>9 m³/t paper produced</td>
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<tr>
<td>Brewing</td>
<td>10−15 m³/m³ beer</td>
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<tr>
<td>Dairy</td>
<td>140 m³/m³ milk</td>
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<tr>
<td>Sugar</td>
<td>8 m³/t sugar</td>
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<tr>
<td>Automotive</td>
<td>450 m³/car (metal production)</td>
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<tr>
<td>Automotive</td>
<td>760 m³/car (tyre production)</td>
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<tr>
<td>Dying</td>
<td>100 m³/t fabric processed</td>
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<td>Soap</td>
<td>2 m³/t soap produced</td>
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<tr>
<td>Power</td>
<td>3 m³/MWh for steam; 60 m³/MWh for cooling</td>
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*aEU figure (Pauly, 2001).  
### Table 1.2 Generalised industrial water quality standards

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<th>5</th>
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<td>Deionised</td>
<td>Purified</td>
<td>Apyrogenic</td>
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<td>-1 to +1</td>
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<td>Silica, mg/l</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td>ND</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TSS, mg/l</td>
<td>&lt; 0.5</td>
<td>&lt; 3</td>
<td>&lt; 3</td>
<td>&lt; 1</td>
<td>&lt; 0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Turbidity, NTU</td>
<td>&lt; 5</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SDI</td>
<td>&lt; 10</td>
<td>&lt; 1</td>
<td>&lt; 1</td>
<td>&lt; 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Particle count, no./ml</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>COD, mg/l</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TOC, mg/l</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Microorganisms, cfu/ml</td>
<td>&lt; 10</td>
<td>&lt; 1</td>
<td>&lt; 1</td>
<td>&lt; 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pyrogens, EU/ml</td>
<td>&lt; 0.25</td>
<td>&lt; 0.25</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

15 years from £1.48 [$2.12] in 1984–85 to £4.90 [$7.94] in 1999–2000 (Water UK, 2000). In more arid regions of the world the figure is higher: between 1980 and 1995 water costs increased by more than an order of magnitude in California (Mannapperema et al., 1993). It is clearly of some cost benefit to acquire a source of water that is commensurate with the quality demanded for its end use, without increasing freshwater demand or discharging waste to sewer. The economic viability of existing municipal effluent reclamation is then very much dependent on the match between the effluent water quality and that demanded by the duty.

A whole industry has built up around the above key concept. Pinch analysis has conventionally been employed to maximise heat transfer efficiency within an industrial process. Water pinch analysis (Baetens, 2002; Section 4.2) is simply pinch analysis based on water quality and volumes, i.e. based solely on mass rather than heat. As such, it is simply an extension of a simple water audit. On the other hand, water pinch analytical tools have not yet reached a level of sophistication where water purification unit processes can be incorporated, since the performance of such processes can rarely be reliably predicted.

Volumes of water used are actually extremely dependent upon the policy of the company or the individual factory regarding water management, and in particular their housekeeping. Housekeeping relates to the way individual unit
### Table 1.3  General water quality requirements for specific duties

<table>
<thead>
<tr>
<th>Use</th>
<th>Industry</th>
<th>Application</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fire fighting</td>
<td>All</td>
<td></td>
<td>1 (natural)</td>
</tr>
<tr>
<td>Irrigation</td>
<td>Agriculture</td>
<td>Drinking water</td>
<td>1 (river)</td>
</tr>
<tr>
<td>Domestic</td>
<td>Offices</td>
<td>Down services</td>
<td>2/3</td>
</tr>
<tr>
<td></td>
<td>Hotels/catering</td>
<td>Laundries</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Healthcare</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Steam raising</td>
<td>Process industries</td>
<td>Heating</td>
<td>3/4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Steam stripping</td>
<td>3/4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High-pressure steam</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Power generation</td>
<td>Turbine drive</td>
<td>8</td>
</tr>
<tr>
<td>Heat transfer</td>
<td>Manufacturing</td>
<td>Closed heating and cooling systems</td>
<td>3/4</td>
</tr>
<tr>
<td></td>
<td>Process industries</td>
<td>Open recirculatory cooling systems</td>
<td>3/4</td>
</tr>
<tr>
<td></td>
<td>Offices</td>
<td>Air conditioning</td>
<td>5</td>
</tr>
<tr>
<td>Process water</td>
<td>Heavy chemicals</td>
<td>Product washing</td>
<td>2/3</td>
</tr>
<tr>
<td></td>
<td>Fine chemicals</td>
<td>Solvent</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Food/soft drinks</td>
<td>Bottle/container washing</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Brewing</td>
<td>Cleaning in place (CIP)</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Pharmaceuticals</td>
<td></td>
<td>6/7</td>
</tr>
<tr>
<td></td>
<td>Metal finishing</td>
<td></td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Photographic</td>
<td></td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Laboratories</td>
<td></td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Semiconductor</td>
<td>Ultrapure water</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>Food/soft drinks</td>
<td>Product quality</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Brewing</td>
<td>Shelf life</td>
<td>2/4</td>
</tr>
<tr>
<td></td>
<td>Pharmaceuticals</td>
<td>Parenterals</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Cosmetics</td>
<td>Lotions/liquids/topicals</td>
<td>5/6</td>
</tr>
</tbody>
</table>

### Table 1.4  Specific water consumption, litres per capita per day, across Europe (Water UK, 2000)

<table>
<thead>
<tr>
<th>Country</th>
<th>Household and small business</th>
<th>Industry and others</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Austria</td>
<td>155</td>
<td>160</td>
<td>100</td>
</tr>
<tr>
<td>Belgium</td>
<td>104</td>
<td>118</td>
<td>59</td>
</tr>
<tr>
<td>Denmark</td>
<td>165</td>
<td>136</td>
<td>96</td>
</tr>
<tr>
<td>Finland</td>
<td>155</td>
<td>155</td>
<td>96</td>
</tr>
<tr>
<td>France</td>
<td>109</td>
<td>151</td>
<td>58</td>
</tr>
<tr>
<td>Germany</td>
<td>139</td>
<td>130</td>
<td>52</td>
</tr>
<tr>
<td>Hungary</td>
<td>110</td>
<td>104</td>
<td>107</td>
</tr>
<tr>
<td>Italy</td>
<td>211</td>
<td>228</td>
<td>69</td>
</tr>
<tr>
<td>Luxembourg</td>
<td>183</td>
<td>172</td>
<td>76</td>
</tr>
<tr>
<td>Netherlands</td>
<td>142</td>
<td>166</td>
<td>37</td>
</tr>
<tr>
<td>Spain</td>
<td>157</td>
<td>145</td>
<td>58</td>
</tr>
<tr>
<td>Sweden</td>
<td>195</td>
<td>188</td>
<td>120</td>
</tr>
<tr>
<td>UK</td>
<td>154</td>
<td></td>
<td>100</td>
</tr>
</tbody>
</table>
operations within the factory are conducted so as, in this instance, to limit the freshwater demand. Actual examples of demand management, reuse and recycling (Table 1.5) demonstrate that the degree of sophistication of the solution to achieve a significant cost benefit depends upon the existing state of water management.

It is apparent from Tables 1.2 and 1.3 that the quality of water demanded by industry varies considerably from one duty to another. There exist certain determinants, such as the Silt Density Index or SDI (Section 2.4.3), which are of critical importance for some applications but are meaningless in others. Moreover, it is generally the case that the volume of water demanded varies inversely with that of its purity across any one industrial sector. For example, according to the figures in Table 1.1, about 20 times as much water—making up around 90% of the total water demand of the plant—is used for cooling in power generation as that required for boiler feed, which demands a "high-purity" water. Such water can only be produced through a combination of adsorptive and membrane separation processes. Water for once-through, non-intrusive cooling, on the other hand, may not be required to meet any specification based on chemical and biological constituents, needing only to be below a certain temperature.

For some industrial processes the quality of the discharged water does not substantially differ from that of the feedwater. Cooling towers, for example, concentrate the water as a result of the evaporative cooling process, but do not add significantly, if at all, to the chemical loading rate, in terms of mass flow rate of solutes, of the effluent. For most industrial sectors, however, there is a significant pollutant load resulting from their activity. As already stated, the large temporal variation in effluent water quality can preclude water recovery and reuse in many cases due to the high cost of treatment to produce water of a reliable quality, particularly by the more established non-barrier technologies where treatment process performance varies with hydraulic and/or pollutant load. On the other hand, membrane processes, which can offer a highly selective barrier to the water being processed, are far more robust to changes in feedwater quality and can provide water of reliably high quality.

### 1.3 Membrane technology

Membrane processes are designed to carry out physical or physicochemical separations. Although most membrane applications are water based, there also exist gas–liquid and gas–gas separation processes, although these are more recent developments and have not yet achieved widespread implementation. In terms of membrane sales, the most important application by far is hemodialysis, as carried out in kidney dialysis machines: almost half of all membrane sales are accounted for by this one application. The development of membrane-based bulk water and wastewater treatment processes, as defined in Table 1.6, is nonetheless significant, since they offer three clear advantages over conventional techniques:
<table>
<thead>
<tr>
<th>Type</th>
<th>Application</th>
<th>Scenario</th>
<th>Solution</th>
<th>Cost benefit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Audit</td>
<td>Cellophane production</td>
<td>750 m$^3$/day of water unaccounted for</td>
<td>Fix leak</td>
<td>£100 000 p.a. saving [£141 580]*</td>
</tr>
<tr>
<td>Audit</td>
<td>Laboratory</td>
<td>Mains water used for single-pass cooling, 12 m$^3$/day</td>
<td>Fit cooler (£2000) [£2831]*</td>
<td>6 month payback</td>
</tr>
<tr>
<td>Housekeeping</td>
<td>Resin factory</td>
<td>Wash water contaminated by spillages from point sources</td>
<td>Fit drip trays to point sources</td>
<td>Effluent COD reduced by 75%.</td>
</tr>
<tr>
<td>Housekeeping</td>
<td>Plating plant</td>
<td>Significant drag out of chemicals from plating bath into rinse bath</td>
<td>Improve drainage around plating bath</td>
<td>£3000 p.a. saving [£4217]*</td>
</tr>
<tr>
<td>Reuse</td>
<td>Plating plant</td>
<td>0.4 m$^3$/h mains water fed to three rinsing baths in parallel, 1.2 m$^3$/h in total</td>
<td>Use counter-current serial flow to all baths</td>
<td>Reduced rinsewater contamination.</td>
</tr>
<tr>
<td>Recycling</td>
<td>Cadmium plating</td>
<td>Zero discharge of cadmium imposed by regulator</td>
<td>Installation of deionisation plant to recover cadmium and water from effluent</td>
<td>£1000 p.a. saving [£1416]*</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>50% water saving, £1500 p.a. [£2124]*</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>12 month payback</td>
</tr>
</tbody>
</table>

* Based on rate of 1.4158.02/06/2002.
Table 1.6  Membrane processes

<table>
<thead>
<tr>
<th>Process</th>
<th>Usual objective</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microfiltration (MF)</td>
<td>Removal of suspended solids, including microorganisms</td>
</tr>
<tr>
<td>Ultrafiltration (UF)</td>
<td>Removal of both large, dissolved solute molecules and suspended colloidal particles</td>
</tr>
<tr>
<td>Nanofiltration (NF)</td>
<td>(Selective) removal of multivalent ions and certain charged or polar molecules</td>
</tr>
<tr>
<td>Reverse osmosis (RO)</td>
<td>Removal of inorganic ions</td>
</tr>
<tr>
<td>Electrodialysis (ED) and dialysis</td>
<td>(Selective) extraction of ions from water and/or concentration of these ions in the waste stream</td>
</tr>
<tr>
<td>Pervaporation (PV)</td>
<td>(Selective) extraction of molecular gas and/or volatile solutes</td>
</tr>
<tr>
<td>Gas transfer (GT)</td>
<td>Transfer of molecular gas into or out of water</td>
</tr>
</tbody>
</table>

1. Separation is achieved without requiring a phase change, and is therefore more energetically efficient than distillation.
2. Little or no accumulation takes place in the process which therefore operates continuously under steady-state condition without necessitating regeneration cycles, unlike adsorptive separation processes.
3. Little or no chemical addition is required, unlike conventional clarification which generally relies on the addition of chemical coagulants and flocculants.

Membrane technology development as a whole began with the first high-performance reverse osmosis membrane produced by in the early 1960s (Loeb and Sourirajan, 1963), which led to the installation of large seawater desalination plant in arid regions of the world. Since that time, growth in the total quantity of membranes sold with reference to the water treatment capacity provided has increased exponentially with time, with growth in microfiltration and ultrafiltration technologies over the last decade of the twentieth century being particularly pronounced (Fig. 1.1). There has been a corresponding exponential decrease in membrane costs with installed membrane plant capacity, as reflected in data from an established hollow fibre microfiltration membrane (Fig. 1.2). The increasing stringency of water quality guidelines and standards being introduced for municipal water treatment, for example the European Union Urban Waste Water Treatment Directive (UWWTD) and the position taken by the UK Drinking Water Inspectorate regarding cryptosporidia, portend continued growth in this sector for membrane technology. Indeed, the EU wastewater treatment membrane market is predicted to increase by an average annual growth rate of 5.9% to $250 million in 2006 (BCC, 2002).

Application of membrane processes within the industrial sector is also widespread and well established in many instances. Reverse osmosis and ultrafiltration, for example, are both widely used within the pharmaceutical industry to remove pyrogens in the provision of water for injection (WFI). These processes are also essential in providing ultrapure water in semiconductor fabrication plant, where ultrafiltration is used for removing colloidal material and reverse osmosis both for primary deionisation (followed by polishing using
Membranes for Industrial Wastewater Recovery and Re-use

Figure 1.1 Total membrane sales with respect to water treatment capacity (by kind permission of Vivendi)

Figure 1.2 Membrane costs vs. installed membrane area ("Memcor" microfiltration plant) (courtesy of Vivendi)

an ion-exchange process) and for polishing the rinse water. Cartridge microfilters are routinely used for the removal of extraneous suspended solids from boiler feedwaters. Although rather less well established, membranes are also used for the treatment of industrial effluents for discharge. Possibly the most recently developed and successfully implemented membrane process for wastewater treatment is the submerged membrane bioreactor, which is applied to both municipal and industrial effluents for the removal of dissolved organic material. Market penetration of this technology has followed much the same trend as that of the more established microfiltration process (Fig. 1.3).

The use of membranes specifically for industrial wastewater reclamation and reuse has generally been constrained by the costs involved. Having said this, advances in membrane technology, and notably significant improvements in its
efficiency and cost effectiveness, greatly increase the competitiveness of recycling over discharge. Existing examples of closed-loop water recovery and reuse, invariably based on membrane technology, suggest that payback periods as low as 18 months are achievable, the timescale obviously being very sensitive to supply and discharge costs. Given the continuing downward pressure on costs (Figs. 1.2 and 1.3) and the increasingly severe stresses on freshwater supplies, it seems likely that membranes will be used ever more widely for water recycling within the industrial sector.

References


Chapter 2

Membrane technology

Simon Judd
School of Water Sciences, Cranfield University
2.1 The membrane

2.1.1 Membrane and membrane process definition

There are a number of definitions of the word “membrane”, which can vary considerably in comprehensiveness and clarity. It is a very widely used term, and can mean anything from a cell wall to damp proofing material. Three definitions, arbitrarily chosen from pertinent technical literature from the last 20 years are provided below:

- “An intervening phase separating two phases and/or acting as an active or passive barrier to the transport of matter between phases” – the European Society of Membrane Science and Technology (now the European Membrane Society).
- “An interphase separating two homogenous phases and affecting the transport of different chemical components in a very specific way” – Prof. Heine Strathmann, former Head of the Department of Membrane Technology, University of Twente.
- “A material through which one type of substance can pass more readily than others, thus presenting the basis of a separation process” – Prof. George Solt, former Director of the School of Water Sciences, Cranfield.

For the purposes of this discussion of membrane technology for wastewater treatment, Solt’s definition can be considered adequate: it is that property of the membrane which permits the separation of components in and/or from water that is of key interest. For many processes the membrane acts to reject the pollutants, which may be suspended or dissolved, and allow the “purified” water through it. In some cases, however, the membrane may act so as to extract pollutants from the wastewater, or else transfer specific components (such as oxygen) into it. Currently employed extractive processes include electrodialysis (ED), dialysis, pervaporation (PV) and gas transfer (GT). In these cases the membrane is employed to allow selective permeation of specific components dissolved in the water. Of far more industrially importance, however, are the filtration processes of reverse osmosis (RO), nanofiltration (NF), ultrafiltration (UF) and microfiltration (MF). In these processes it is the bulk water that passes through the membrane under an applied pressure, leaving the pollutants in concentrated form on the unpermeated side of the membrane. If hemodialytic applications are ignored, then the pressure-driven filtration processes account for around 75% of the remaining membrane sales, almost all of these due to aqueous separation applications in municipal and industrial water treatment.

2.1.2 Membrane structure

Although membrane materials vary vastly according to chemical composition and process type, the principal objectives in membrane manufacture are always the same. An ideal material will:
Membrane technology

- have reasonable mechanical strength,
- maintain a high throughput, and
- be selective for the desired permeate constituent.

These last two parameters are mutually counteractive, since a high degree of selectivity is normally only achievable using a membrane having small pores and thus an inherently high hydraulic resistance (or low permeability). The permeability also increases with increasing density of pores, and the overall membrane resistance is directly proportional to its thickness (in accordance with Darcy's law). Finally, selectivity will be compromised by a broad pore size distribution. An optimum physical structure for any membrane material is thus:

- a thin layer of material,
- a narrow range of pore size, and
- a high porosity.

Membrane materials can be categorised as either dense or porous, and by the mechanism by which separation is actually achieved (Table 2.1). Separation by dense membranes relies to some extent on physicochemical interactions between the permeating components and the membrane material, and relate to separation processes having the highest selectivity (Fig. 2.1). Porous membranes, on the other hand, achieve separation mechanically by size exclusion (i.e. sieving), where the rejected material may be either dissolved or suspended depending on its size relative to that of the pore. Since some membranes exhibit properties that can be associated with more than one process type, the boundaries between the adjacent membrane processes in Fig. 2.1 are somewhat nebulous. For example, IUPAC (1985) state that the upper and lower boundary limits for mesopores, as are characteristic of a UF membrane, are 2 and 50 nm. According to Kesting (1989), however, these boundaries are at 1 and 20 nm, respectively.

Whilst microfiltration membranes are assigned a characteristic pore size, the exact value of which is dependent on the method of measurement, ultrafiltration membranes are mostly rated on the basis of the size of the smallest molecule the membrane can be expected to reject. This is routinely expressed as the molecular weight cut-off (MWCO) in daltons (i.e. grams per mole). As the precise relationship between MWCO and pore size is obviously dependent on the physical and chemical nature of the solute molecule, precise cross-referencing is impossible. The actual pore size of nanofiltration and reverse osmosis membranes is of little practical consequence, since there are other mechanisms more dominant than simple sieving that determine membrane performance. The purification performance of these membranes can only be rated according to their actual demonstrated permselectivity, i.e. the extent of the rejection of key contaminants by the membrane, under some defined set of conditions. Nanofiltration membranes, which have a charge rejection component, are generally designed to be selective for multivalent rather than univalent ions. Reverse osmosis membranes are designed to reject all species other than water.
Table 2.1 Dense and porous membranes for water treatment (adapted from Stephenson et al., 2000)

<table>
<thead>
<tr>
<th>Dense</th>
<th>Porous</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Membrane separation processes</strong></td>
<td><strong>Ultrafiltration (UF)</strong></td>
</tr>
<tr>
<td>Reverse osmosis (RO)</td>
<td>Separation by sieving through mesopores (2–50 nm)*a.</td>
</tr>
<tr>
<td>Separation achieved by virtue of differing solubility and diffusion rates of water (solvent) and solutes in water.</td>
<td>Microfiltration (MF)</td>
</tr>
<tr>
<td>Electrodialysis (ED)</td>
<td>Separation of suspended solids from water by sieving through macropores (&gt; 50 nm)*a.</td>
</tr>
<tr>
<td>Separation achieved by virtue of differing ionic size, charge and charge density of solute ions, using ion-exchange membranes.</td>
<td>Gas transfer (GT)</td>
</tr>
<tr>
<td>Pervaporation (PV)</td>
<td>Gas transferred under a partial pressure gradient into or out of water in molecular form.</td>
</tr>
<tr>
<td>Same mechanism as RO but with the (volatile) solute partially vaporised in the membrane by partially vacuumating the permeate.</td>
<td></td>
</tr>
<tr>
<td>Nanofiltration (NF)</td>
<td></td>
</tr>
<tr>
<td>Formerly called leaky reverse osmosis. Separation achieved through combination of charge rejection, solubility-diffusion and sieving through micropores (&lt; 2 nm).</td>
<td></td>
</tr>
<tr>
<td>Membrane materials</td>
<td>Both polymeric and inorganic materials available.</td>
</tr>
<tr>
<td>Limited to polymeric materials.</td>
<td></td>
</tr>
</tbody>
</table>

*a IUPAC (1985).

although they are unable to offer a significant barrier to dissolved gases and certain low-molecular-weight organic molecules.

Membranes may also be categorised according to the material composition, which is either organic (polymeric) or inorganic (ceramic or metallic), or on the basis of their physical structure, i.e. their morphology. The membrane morphology is dependent on the exact nature of the material and/or the way in which it is processed. In general, however, membranes employed in pressure-driven processes tend to be anisotropic: they have symmetry in a single direction, and hence are often referred to as asymmetric, such that their pore size varies with membrane depth (Fig. 2.2). This arises out of the requirement for a thin permselective layer (or skin) to minimise the hydraulic resistance, the porous support providing minimal resistance and acting purely to provide the necessary mechanical strength. For integral (i.e. single, non-composite material) asymmetric polymeric membranes, such as the one shown in Fig. 2.2a, the skin is normally 2–5 μm in thickness.

Flat sheet reverse osmosis membranes have an additional ultrathin active layer, less than 0.5 μm in thickness, attached to the anisotropic substrate to produce a thin-film composite (TFC) membrane (Fig. 2.3). This ultrathin layer provides the required permselectivity, rejecting all charged species and permitting only the passage of water and small organic molecules. On the other hand, ion exchange membranes, which are also dense by definition, are essentially homogeneous. These membranes comprise a three-dimensional array of fixed ionogenic sites (i.e. functional groups capable of dissociating to form charged species) which facilitate transport of either cations or anions.
Figure 2.1 Membrane separation processes overview (from Stephenson et al., 2000, reproduced with permission)

Figure 2.2 Anisotropic UF membranes: (a) polymeric (thickness of "skin" indicated), and (b) ceramic (by kind permission of Ionpure)

through the membrane. Certain porous membranes are also homogeneous, but these tend to have a correspondingly low permeability.

2.1.3 Membrane materials and their manufacture

Membrane manufacture ostensibly concerns the production of a permeable material at a reasonable cost. The membrane cost is dependent on not only the
raw material but also on the ease with which pores of the desired size or size distribution can be introduced. This can vary considerably from one material to the next according to the method used and the corresponding precision of the pore size distribution (or degree of isoporosity).

The range of available membrane materials employed in water and wastewater treatment is very broad, and they vary rather more widely in chemical composition than in bulk morphology. Membranes can be produced by stretching, sub-atomic particle bombardment combined with etching and, in the case of ceramic materials, sintering (Table 2.2). These membranes are formed by the pressing and sintering of fine powders onto a pre-prepared porous support, which can produce pore diameters down to around 2 μm, followed by sol–gel processes to produce successive deposits of progressively finer porous layers down to a minimum pore size of around 3 nm. Ceramic nanofiltration membranes, however, have only recently arrived in the marketplace. The production process tends to be very expensive if a highly selective membrane layer of even thickness and narrow pore size distribution is to be produced. The cost of microfiltration or ultrafiltration membranes derived from titanium and/or zirconium usually exceeds $1000 per m², although there appears to be some progress in producing low-cost ceramic membranes. At the opposite end of the spectrum are simple, homogenous polymeric membranes produced by extrusion (stretching) of partially crystalline sheets perpendicular to the orientation of the crystallites, possibly with the assistance of a fibrulating agent. Such materials cost less than $1 per m² to produce, since the process can be made continuous, and the cost of the membrane module is then determined almost entirely by the module fabrication cost. However, such membranes are limited in their permeability, isoporosity and mechanical strength.

Supported liquid membranes (SLMs) have not yet been produced on a commercial scale, and currently appear to be more appropriate to gas separations where the improved mass transport of gases through liquids over that attainable in solids becomes important. SLMs currently being developed for
Membrane technology

Table 2.2 Membrane materials by type

<table>
<thead>
<tr>
<th>Membrane</th>
<th>Manufacturing procedure</th>
<th>Applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ceramic</td>
<td>Pressing, sintering of fine powders followed by sol-gel coating</td>
<td>MF, UF, Aggressive and/or highly fouling media</td>
</tr>
<tr>
<td>Stretched polymers</td>
<td>Stretching of partially crystalline foil</td>
<td>MF. Aggressive media, sterile filtration, medical technology</td>
</tr>
<tr>
<td>Track-etched polymers</td>
<td>Radiation followed by acid etching</td>
<td>MF (polycarbonate (PC) or polyethylene terephthalate (PET) materials). Analytical and medical chemistry, sterile filtration</td>
</tr>
<tr>
<td>Supported liquid</td>
<td>Formation of liquid film in inert polymer matrix</td>
<td>Gas separations, carrier-mediated transport</td>
</tr>
<tr>
<td>Integral asymmetric, microporous</td>
<td>Phase inversion</td>
<td>MF, UF, NF, GT</td>
</tr>
<tr>
<td>Composite asymmetric, microporous</td>
<td>Application of thin film to integral asymmetric microporous membrane to produce TFC</td>
<td>NF, RO, PV</td>
</tr>
<tr>
<td>Ion exchange</td>
<td>Functionalisation of polymer material</td>
<td></td>
</tr>
</tbody>
</table>

liquid separations usually comprise a high-viscosity, hydrophobic liquid immobilised in a polymer matrix. The supported liquid may contain a carrier, a component that reacts chemically and reversibly with the desired component in the liquid mixture and thereby assists its transport through the membrane.

By far the most significant of the polymeric membranes applied to the water industry are those produced by phase inversion (Table 2.3), a fabrication process which produces an anisotropic material which may then be used as a substrate for a TFC membrane. The ultrathin surface layer of a TFC RO or NF membrane is invariably either a polyamide (usually aromatic) or some co-polymer blend based on polyamide. The permselectivity and throughput of such membranes is then critically dependent upon the precise polymers selected and the fabrication method used. For example, Filmtec (part of the Dow Chemical Company) produce both a reverse osmosis membrane (the FT-30) and a series of nanofiltration membrane materials (NF45, NF70 and NF90) which are all based on aromatic polyamide, yet the permeability and salt rejection capability of these membranes varies considerably.

The phase inversion process involves dissolving the polymer in a suitable solvent and then casting it in a film, less than 1 mm thick, and then adding another liquid to precipitate the polymer. The membrane skin forms at the interface between the solvent and the second liquid, in which the membrane is only sparingly soluble. Careful choice of the solvent and non-solvent liquids, concentration of the polymer, temperature and reaction times can produce the desired physical membrane characteristics.

Most polymeric materials are resistant to moderate pH swings, the principal exception being cellulose acetate, but most have limited resistance to organic solvents. Only PTFE and PVDF can be considered highly stable in this regard. The main limitation of the more robust polymeric materials, in terms of resistance to
### Table 2.3 Commercially available phase inversion polymeric membrane materials

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Process</th>
</tr>
</thead>
<tbody>
<tr>
<td>CA</td>
<td>Chlorine resistant, inexpensive, more fouling resistant than PA</td>
<td>Susceptible to alkaline hydrolysis at pH &gt; 6, susceptible to biodegradation, limited thermal and chemical stability, limited permselectivity (~95% rejection)</td>
<td>RO, NF, UF</td>
</tr>
<tr>
<td>PA</td>
<td>More all-round stability than CA, more permselectivity than CA, high resistance to hydrolysis</td>
<td>Very limited chlorine tolerance (&lt; 0.1 mg l(^{-1}))</td>
<td>RO, NF</td>
</tr>
<tr>
<td>PAN</td>
<td>High resistance to hydrolysis, high resistance to oxidation</td>
<td>Hydrophobic, requires copolymers to make less brittle</td>
<td>UF, RO substr.</td>
</tr>
<tr>
<td>PSU, PES</td>
<td>Very good all-round stability, mechanically strong</td>
<td>Hydrophobic</td>
<td>UF, RO substr.</td>
</tr>
<tr>
<td>PVDF, PTFE</td>
<td>Extremely high chemical stability, high thermal stability</td>
<td>Highly hydrophobic, limited mechanical stability, limited intrinsic permeability, expensive</td>
<td>UF, MF</td>
</tr>
<tr>
<td>PEI</td>
<td>High chemical stability, very high thermal stability, mechanically strong</td>
<td>Hydrophobic, less solvent resistant than PVDF, poorer alkaline stability than PSU or PAN</td>
<td>UF, RO substr.</td>
</tr>
<tr>
<td>PP</td>
<td>Inexpensive</td>
<td>Hydrophobic</td>
<td>UF, MF</td>
</tr>
</tbody>
</table>

- CA, cellulose acetate (predominantly di- or tri-acetate); PA, polyamide (aromatic); PAN, polyacrylonitrile; PSU, polysulphone; PES, poly(ether sulphone); PVDF, poly(vinyldene fluoride); PTFE, poly(tetrafluoroethylene); PEI, polyetherimide; PP, polypropylene.
- Most usual application in bold type.

Oxidative attack and hydrolytic degradation promoted by extremes of pH, is their hydrophobicity. This makes them very susceptible to non-specific adsorption of hydrophobic contaminants in the water, which reduce the permeability of the membrane. This phenomenon, termed fouling, imposes a very significant constraint on the operation of all membrane processes. Much developmental work has been conducted in producing chemically and mechanically robust membranes with hydrophilic surfaces that are therefore less susceptible to fouling by common natural hydrophobic materials such as proteins.

Surface modification can take place by four recognised routes:

- chemical oxidation,
- organic chemical reaction,
- plasma treatment, and
- grafting.

Chemical oxidation, using standard oxidative chemicals such as chromic or nitric acid, converts surface groups such as alkenes and esters to hydroxyl and
carboxylate groups, respectively, which are more hydrophilic and normally negatively charged. Oxidative chemicals can also be used to promote grafting of hydrophilic monomers onto the membrane surface. Plasma treatment involves the generation of a plasma by continuous electrical discharge in a gas containing oxygen or water (to produce oxygen-containing functional groups), or nitrogen, ammonia or amines (to produce nitrogen-containing functional groups). Plasma treatment offers the advantages of being fairly controllable, since the reaction conditions can be adjusted to produce the required degree of reaction, and demands no hazardous liquids.

Chemical reaction and grafting add an organic layer to the surface of the polymer, rather than functionalising it in the way that the above oxidative reactions do. The surface can also be modified by specific organic chemical reactions which rely on the chemical reactivity of the polymer. Polysulphone, for example, will react with chloroether to produce the ortho-chloromethylated polymer (Guiver et al., 1993). Polymer grafting can be promoted chemically or thermally but more usually involves irradiation in the presence of a suitable initiator and, invariably, vinyl or acrylic monomers.

Ion exchange membranes, whose most important property is permeability to counter-ions and impermeability to co-ions, can be produced either by functionalisation of a membrane, as mentioned above, or by immobilising powdered ion exchange resins in an inert resin matrix. Functionalisation produces a homogenous material that, as with many membranes, requires fixing on a suitable material to give it mechanical strength. The powder immobilisation method is relatively inexpensive but the materials produced are heterogeneous, less selective for counter-ions and have inferior mass transport properties than the equivalent more expensive homogeneous materials.

Recent developments: membrane materials
The search for new membrane materials with superior properties with respect to permeability and resistance to chemical, thermal and biological attack is ongoing. Of greatest interest is fouling resistance, and this demands the development of membranes with a low affinity to pre-identified foulants in the feedstock. The ease with which this can be achieved is to a large extent dependent on the heterogeneity of the feedwater matrix, since the membrane surface cannot be modified so as to be able to repel contaminants of different charges and hydrophobicities. In general natural foulants tend to carry a negative charge, which then demands that the membrane also carries a negative charge to repel the foulant. It should be pointed out, however, that some filters are surface modified to carry a positive charge (e.g. the Pall "Posidyne" nylon-based filter) to attract negatively charged suspended materials, but these filters are actually depth filters designed to retain particles rather than reject them.

One application that has received a great deal of interest is the removal of natural organic matter (NOM) from upland surface waters for potable water production using nanofiltration membranes. NOM contains a number of different fractions of which humic acids are one. Humic acids, which contain phenolic and carboxylate functional groups, tend to be negatively charged in
neutral solutions, such that a highly negatively charged membrane would be expected to repel these contaminants allowing operation at higher fluxes than would normally be attainable without fouling. This has focused attention upon highly negatively charged membranes and, in particular, blends of sulphonated poly(ether ether ketone) (SPEEK) or sulphonated poly(ether ether sulphone) (SPEES) with polysulphone or polyethersulphone, respectively (Bowen et al., 2001; Knoell et al., 1999). These membranes are likely to be commercialised within the next five years.

Another important constraint on the application of membranes to industrial process waters is their resistance both to extreme pH conditions and, more especially, key organic solvents. A few porous membrane materials, notably PTFE and PVDF, are intrinsically resistant to attack from both chlorinated and hydroxylated or carboxylated organic solvents, whereas PSU, PES and PEI will withstand some organic solvents but not chlorinated reagents like chloroform. Recently, a range of organic nanofiltration membranes (SelRO\textsuperscript{K}, commercialised by Koch) have become available that can apparently be employed in pure, non-chlorinated organic solvents. These membranes are probably a modified cross-linked PAN material. Details of other solvent-resistant UF and NF membranes either recently available or under development are provided by Nunes and Peinemann (2001), and this information is summarised in Table 2.4.

So far as ceramic filters are concerned, there is a general drive to increase their competitiveness by reducing costs to allow them to be sold at less than $1000 per m\textsuperscript{2}, although it is still the case that such a price is only attainable for large bulk orders. However, very recently the opportunities offered by extrusion of certain ceramic materials have started to be explored, for example by Ceramem, allowing the production of membrane materials of much lower cost (Section 2.1.4). Perhaps the most intriguing of the ceramic filters, however, is the Anopore filter. This is produced by electro-oxidation of aluminium to produce an almost perfect honeycomb structure with an extremely narrow pore size distribution and a surface porosity of up to 50%. This porosity value compares well with the similarly homoporous Nuclepore membranes, produced by neutron bombardment followed by track-etching of polyethylene terephthalate.

<table>
<thead>
<tr>
<th>Material</th>
<th>Application</th>
<th>Supplier(s) or developer(s)</th>
<th>Stability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrophilic</td>
<td>NF</td>
<td>Koch SelRO\textsuperscript{K}</td>
<td>DMF, NMP, DMSO resistant: 0–14 pH, 70°C</td>
</tr>
<tr>
<td>PAN</td>
<td>NF</td>
<td>Desal D-series, Osmonics</td>
<td>35% H\textsubscript{2}SO\textsubscript{4}, 25% H\textsubscript{3}PO\textsubscript{4}</td>
</tr>
<tr>
<td></td>
<td>NF</td>
<td>Somicon/Nitto</td>
<td>15% NaOH, 60°C, pH &lt; 1</td>
</tr>
<tr>
<td>PVDF/Cellul.</td>
<td>UF/NF</td>
<td>Dow Danmark Etrna</td>
<td>pH 1–12</td>
</tr>
<tr>
<td>PLL</td>
<td>UF</td>
<td>Nitto, Bend Research</td>
<td>–</td>
</tr>
<tr>
<td>PEEK</td>
<td>UF</td>
<td>Dow, Ionics</td>
<td>–</td>
</tr>
<tr>
<td>PAEK/PEI</td>
<td>UF</td>
<td>Bend Research</td>
<td>–</td>
</tr>
</tbody>
</table>
which is no more than 14% porous. Unfortunately, the membrane thickness of 60 μm makes the hydraulic resistance of the Anopore membranes relatively high and, more crucially, they can currently only be produced in small batches as disc filters for use in the laboratory.

2.1.4 Membrane configurations

A membrane is only useful if it takes a form which allows water or pollutants to pass through it. The configuration of the membrane, i.e. its geometry and the way it is mounted and oriented in relation to the flow of water, is crucial in determining the overall process performance. Other practical considerations concern the way in which the membrane elements, i.e. the individual discrete membrane units themselves, are housed to produce modules, the complete vessels through which the water flows. The optimum membrane configuration is one that has the following characteristics:

(a) a high membrane area to module bulk volume ratio
(b) a high degree of turbulence for mass transfer promotion on the feed side
(c) a low energy expenditure per unit product water volume
(d) a low cost per unit membrane area
(e) a design that facilitates cleaning
(f) a design that permits modularisation.

All membrane module designs, by definition, permit modularisation (f), and this presents one of the attractive features of membrane processes per se. However, some of the remaining listed characteristics are mutually exclusive. For example, promoting turbulence (b) results in an increase in the energy expenditure (c). Direct mechanical cleaning of the membrane (e) is only possible on comparatively low area:volume units (a) where the membrane is accessible. Such module designs inevitably increase the total cost per unit membrane area (d). Finally, it is not possible to produce a high membrane area to module bulk volume ratio without producing a unit having narrow feed channels, which will then adversely affect turbulence promotion.

There are five principal configurations currently employed in membrane processes, which all have various practical benefits and limitations (Table 2.5). The configurations are based on either a planar or cylindrical geometry and comprise:

- Pleated filter cartridge
- Plate-and-frame
- Spiral wound
- Tubular
- Hollow fibre

Of the flat plate geometries the lowest cost is the pleated filter cartridge (Fig. 2.4), used exclusively in microfiltration and designed as a relatively inexpensive
disposable unit to be employed for polishing relatively pure waters. Pleated media provide a high area to volume ratio and also ensure the minimum number of sealing operations, i.e. a side seal and two end seals. The construction is such that the loss of filter area at the folds of the pleat is avoided by using a suitably designed membrane separator. The configuration is suitable for both organic polymeric media and porous metal media, the latter having a minimum pore size of around 2 μm. For these filters, and for other membrane modules applied to pure waters, key design parameters are integrity testability, sterilisability and, most importantly, propensity to leach contaminants from the module or unload retained particles from the filter medium.

Plate-and-frame modules comprise rectangular or (pseudo) circular flat sheet membranes with separators and/or support plates. The modules can be pressure sealed, such as with electrodialysis stacks, or be designed as cassettes. Cassettes allow membrane elements to be inserted and removed individually without the necessity of partially dismantling the module. They are, however, limited to low-pressure operation, and hence to microfiltration or ultrafiltration processes. An example is the Kubota membrane, employed exclusively in a membrane bioreactor (Fig. 2.5). This technology operates at transmembrane pressures as low as 0.05 bar, and therefore does not demand as robust a construction as a pressure-sealed unit would provide. The membrane element comprises two rectangular sheets, 0.4 m x 0.8 m, heat sealed at the perimeters to a plastic frame. The elements are inserted into a cassette holding up to 150 elements, with the permeate collected from inside each one individually.

An example of a circular pressure-sealed module is the Rochem reverse osmosis module (Fig. 2.6). This module employs support plates of 200–400 mm diameter and 5 mm thick at the outside edge. The plate surfaces are covered with tiny protuberances that promote turbulence as the water flows over them. Sandwiched between adjacent plates is a membrane element comprising two membrane sheets, heat sealed at the edges and separated by a thin plastic mesh. The feed is introduced at the top of the stack and flows radially outward at the top

---

**Table 2.5 Membrane configurations**

<table>
<thead>
<tr>
<th>Configuration</th>
<th>Area/vol. ratio (m²/m³)</th>
<th>Cost</th>
<th>Turbulence promotion</th>
<th>Backflushable?</th>
<th>Application*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pleated cartridge</td>
<td>500–1500</td>
<td>Very low</td>
<td>Very poor</td>
<td>No</td>
<td>DEME, low TSS waters</td>
</tr>
<tr>
<td>Plate-and-frame</td>
<td>100–300b</td>
<td>High</td>
<td>Fair</td>
<td>No</td>
<td>ED, UF, RO</td>
</tr>
<tr>
<td>Spiral wound</td>
<td>800–1200</td>
<td>Low</td>
<td>Poor</td>
<td>No</td>
<td>RO, NF, UF</td>
</tr>
<tr>
<td>Tubular</td>
<td>150–300</td>
<td>Very high</td>
<td>Very good</td>
<td>No</td>
<td>CFMF, high TSS waters</td>
</tr>
<tr>
<td>Capillary tubec</td>
<td>1500–5000</td>
<td>Low</td>
<td>Good</td>
<td>Yes</td>
<td>UF</td>
</tr>
<tr>
<td>Hollow fibred</td>
<td>10 000–20 000</td>
<td>Very low</td>
<td>Very poor</td>
<td>Yes</td>
<td>MF, RO</td>
</tr>
</tbody>
</table>

* Most important application in bold type; DE = dead-end, CF = cross-flow.

b Can be 2/m³ for a cassette.
c Capillary tube used in UF: water flows from inside to outside the tubes.
d Hollow fibre used in MF and RO: water flows from outside to inside the tubes.
of the membrane element and radially inwards at the bottom whilst a fraction of
the flow permeates the membrane to form the product water inside the element.
The retentate, i.e. the rejected concentrated water from a single elemental stage, is
then transferred to the next element. There may be hundreds of elements in a single
stack, producing a very long retentate flow path and allowing commensurately
high conversions, i.e. percentage formation of product water from feed water.
This module can also be configured for each element to be arranged in series
rather than in parallel, giving very short retentate flow paths, which reduce
concentration polarisation-exacerbated fouling (Section 2.2.3) to some extent.

Examples of rectangular pressure-sealed plate-and-frame modules are the
Rhodia-Orelis PLAEIDE UF/MF module (Fig. 2.7) and electrodialysis stacks (Fig.
2.8). The latter are all based on a plate-and-frame design since the process relies
on transport of ions through the membranes under the action of an electric field and this configuration presents the simplest means of exposing a large membrane area to an orthogonal electric field. Electrodialysis is simply a combination of electrolysis and ion-selective permeation. If a potential difference (voltage) is applied across a solution of ions they are transported through the solution by electromigration, giving rise to the flow of a current. If ion-selective membranes are then placed in the solution then only the counter-ions are allowed to pass through the membrane via the oppositely charged pores in membrane material. The placement of alternate cation and anion selective membranes (Fig. 2.8) produces compartments or cells in which the salts are alternately concentrated and demineralised by the action of the applied voltage and the resulting flow of current.

The nature of the ED process makes for a module that is somewhat complex in design, and is thus comparatively expensive, and which has to be hand
Membrane technology  27

Figure 2.8  Electrodialysis stack (by kind permission of Ionics)

Assembled. By the same token, the design allows both membrane sheets and the spacers that separate them to be removed and replaced on an individual basis. Indeed, ED suppliers are able to tailor the membrane spacer to the feedwater specification and the duty to which the technology is being put, allowing a degree of flexibility that other membrane module designs are unable to provide (Section 2.4.2). Ceramic rectangular pressure-sealed plate-and-frame modules also exist, an example being the Keram module by TAMI.

Spirally wound membranes (Fig. 2.9) have the advantage of simple and robust construction whilst providing a reasonable membrane area per unit volume, and find use in reverse osmosis and nanofiltration plant. As with many membrane modules, the membrane element comprises two membranes sandwiched together, in this case forming a rectangular “bag” sealed on three sides. The open side is presented to a central perforated tube which collects the permeate product from inside the membrane element. The membrane element is rolled up to produce a spiral which forms a cylindrical shape, one end of which is presented to the feedwater. The fluid path length then equates to the length of the cylinder, and spacers are used to maintain separation of the membrane leaves both in the retentate, where the spacers also serve the important task of turbulence promotion, and in the permeate channels. A number of membrane elements, up to seven for very large plant, can make up a single module by
linking the retentate streams in series and placing the module in an appropriate pressure vessel.

Tubular modules provide a modest surface area to volume ratio, and thus the highest cost per unit area of all cylindrical membrane geometries, but also provide potentially the greatest turbulence promotion and the best access to the membrane surface. Both these factors tend to favour this configuration for highly fouling matrices. If the tube diameter is sufficiently large it is possible to mechanically clean the membranes in place, as is carried out with the FYNE\textsuperscript{N} nanofiltration process developed by PCI Membranes. This process uses foam balls to wipe the inner surface of the membrane tubes once fouled, thus minimising the use of chemicals for this duty. Commercially available modules are mainly multi-channel (Fig. 2.10), with mechanical support required for the polymeric tubes. The ceramic tubes tend to have a lower packing density due to their monolith construction: the tubes comprise cylindrical holes bored into a solid ceramic support. The brittle nature of the ceramic material imposes a lower limit on the tube wall thickness, tending to increase the overall membrane hydraulic resistance over that of a comparable polymeric membrane.

The inherent radial mechanical strength of a tube increases with decreasing diameter, such that at a certain diameter the tube becomes self-supporting. This is the case for capillary tubes and hollow fibres. The highest packing densities are obtained from hollow fibre membrane modules (Fig. 2.11). These can have inner diameters as small as 10 µm for particularly fine fibres (hence hollow fine fibres, or HFF), which therefore give the highest packing densities and so the lowest cost. The module is of a very simple construction. The bundle of fibres is simply bent double and the ends potted and exposed at one end of the tube into which they are inserted. Hollow fibre modules operate out-to-in, such that the skin is formed on the outer membrane surface. Capillary fibres modules, which are slightly larger filaments, are operated in-to-out with the skin therefore formed on
Membrane technology

Figure 2.10 Tubular UF modules: (a) polymeric (PCI Membranes), and (b) ceramic (Kerasep®, by Rhodia Orelis)

Figure 2.11 Hollow fibre module

the internal surface. This allows more control of the hydrodynamics, but the operational pressure is then limited by the mechanical strength of the fibres, which relates to the wall thickness. However, both capillary fibres and hollow fibres can be backflushed (i.e. the flow reversed to wash the foulant from the membrane surface). This is particularly crucial for hollow fibres, where the lack of hydrodynamic control is such that fouling is inevitable and a periodic backflush cycle is essential to maintain a reasonable permeability. Backflushing can be enhanced by simultaneous air sparging, as is the case with the Memcor system.

Recent developments: module configurations

There has been very little development in terms of novel membrane geometries over the last few years: virtually all membrane elements are configured as either planar or cylindrical. A noteworthy development is the newer 27-channel Kerasep® ceramic membrane element produced by Rhodia Orelis, which provides almost 40% more membrane area per unit volume than their conventional cylindrical 19-channel tubular monolith. Each channel is a 120° segment, such that the walls dividing a cluster of three channels are planar. This allows a greater membrane area without compromising mechanical strength. There has also been a tendency to try and produce less expensive modules either by using alternative materials or designs that are less complicated in
manufacture. Hence, some companies have started producing cassettes alongside their pressure-sealed module products.

One noteworthy recent development concerns ceramic membranes. As already stated, the onus is on suppliers of these membranes to reduce costs to make them economically competitive with the much less expensive polymeric materials. It appears that this might be achievable by extruding ceramic membrane substrates to produce channels of rectangular geometry (Fig. 2.12). The cost of these membranes, produced from silicon carbide substrate monolith coated with titania, alumina or zirconia, can be as low as $100 per m² for narrow-membrane channels (2 mm).

Another fairly recent development is the use of aeration combined with submerged membrane systems. Submerged (or immersed) membrane systems, where large-area membrane filtration modules are submerged in a tank and the permeate removed at very low transmembrane pressures, were originally developed for membrane bioreactors (MBRs) by companies like Kubota and Zenon in the late 1980s as an alternative to the more conventional sidestream (or external) filtration systems. MBRs are an example of a hybrid process, in this case combining the activated sludge aerobic biotreatment process with membrane filtration (Stephenson et al., 2000). The high fouling propensity of the sludge liquor being filtered, which can have a suspended concentration of up to 20 g l⁻¹, demands a high degree of turbulence promotion to enhance the flux. In the submerged system turbulence promotion is provided by coarse bubble aeration. It has been shown by a number of researchers (Ghosh and Cui, 1999; Mercier et al., 1997; Cabassud et al., 1997) that air–water two-phase flow can substantially enhance the membrane flux over that attained by single-phase

Figure 2.12  Ceramem silicon carbide monolith substrate (reproduced with permission)
pumped liquid flow. Such has been the success of the application of coarse aeration to submerged MBR systems in municipal and industrial effluent treatment that submerged aerated systems are now starting to be applied to largely abiotic pure water systems.

Another significant innovation is the mechanically enhanced, high-shear processes. These actually employ conventional stacked disc modules, but the module is made considerably more effective by actually moving the membrane in relation to the flowing liquid. These products, which include vibratory shear enhanced processing (VSEP™, marketed by New Logic, the similar Vibrating Membrane Filter, or VMF™, marketed by Pall) and the Valmet-Flootek CR™ membrane processes (Fig. 2.13), enhance the flux through a UF or MF membrane by imposing a shear at or close to its surface. Since the throughput of all membrane processes operating under steady-state conditions is ultimately dictated by shear rate (Section 2.2.2), very substantial increases in throughput are attainable through this modification. These systems are generally considered to be too energy intensive for application to anything other than high-added-value process waters, but all have been employed for resource recovery.

The systems work in different ways in achieving the high shear rates desired. The VSEP process imposes an oscillatory motion on the whole module, moving it just a few degrees around the axis at a rate of around 45 s⁻¹. It is claimed that this imposes a shear rate at the membrane of 150 000 s⁻¹, about ten times that attainable from conventional cross-flow filtration (Section 2.2.1). Although usually applied to a stacked disc filter, because the V-Sep system involves the mechanical agitation of the entire membrane module it can also be applied to other membrane configurations. The VMF system employs a similar high-frequency vibration. The CR filter employs rotors above each membrane element in the stacked disc, and is thus integral to the membrane module. The angular

![Figure 2.13](image_url)  
*Figure 2.13* High-shear processes: (a) VSEP™ resonating drive system (on which the membrane module sits), and (b) CR filter
velocity of the rotor is around 10–15 m s⁻¹, depending on the diameter of the membrane cell. Both the VSEP and CR ultrafiltration processes have been successfully applied to liquors of high suspended solids content (Sections 3.2.5 and 5.6), achieving concentrate streams in excess of 5% solids in some cases.

Other turbulence promotion modifications to modules are still largely at the developmental stage. These include intermittent jets, in which the feed is pumped coaxially through a membrane tube at fixed intervals through a nozzle. The abrupt change in velocity produces a toroidal vortex, increasing the flux by up to 2.5 times for granular suspensions such as bentonite clay (Arroyo and Fonade, 1993). Pulsed flow has also received much attention (Gupta et al., 1985, 1992; Rodgers and Sparks, 1993; Bertram et al., 1993). In this mode, pulses of flow are generated in the feed or permeate channel, again creating large temporal changes in the velocity gradient. A simpler alternative for turbulence promotion is the use of simple inserts in tubes. This is also not practised commercially, despite the many publications in this area, as reviewed by Gupta et al. (1995), Belfort et al. (1994) and others.

Finally, a process that has significantly extended the capability of electrodialysis is the electrodeionisation (EDI), also called continuous deionisation (CDI), process commercialised by Vivendi Ionpure and Ionics. In this process the diluate cells of the electrodialysis stack are filled with ion exchange resin beads. The resin effectively aids the transport of ions from the diluate to the concentrate cells by providing a conducting pathway. This results in extremely effective removal of all charged species in the diluate cell, producing a product water of a quality comparable to that from a twin-bed deionisation process but offering the advantage of being continuous.

2.2 The process fundamentals

2.2.1 Process performance definitions

Flux
The key elements of any membrane process are the influence of the following parameters on the overall permeate flux:

- the membrane resistance,
- the operational driving force per unit membrane area,
- the hydrodynamic conditions at the membrane–liquid interface, and
- the fouling and subsequent cleaning of the membrane surface.

The flux is the quantity of material passing through a unit area of membrane per unit time. This means that it takes SI units of m³ m⁻² s⁻¹, or simply m s⁻¹, and is occasionally referred to as the permeate velocity. Other non-SI units used are l m⁻² h⁻¹ (or “LMH”) and m³ per day, which tend to give more accessible
numbers: membranes generally operate at fluxes between 10 and 1000 LMH. The flux relates directly to the driving force (Section 2.2.2) and the total resistance offered by the membrane and the interfacial region adjacent to it.

Conversion
In membrane processes there are three possible streams: a feed, a retentate and a permeate stream. The retentate stream is unpermeated product. If there is no retentate stream then operation is termed dead-end or full-flow (Fig. 2.14a). Such operation is normally restricted to either low-solids water, as for cartridge filtration of boiler feedwater or ultrafiltration for apyrogenic pure water production, or cyclic operation with frequent backwashing, such for most microfiltration and ultrafiltration membrane plant for municipal water treatment. For waters having a significant solids loading and/or membranes of limited permeability (dense membranes), it is not desirable to try and convert all of the feed to permeate product in a single passage through a module. In such cases, cross-flow operation is employed (Fig. 2.14b) whereby some of the feedwater is collected as a concentrate (or retentate) stream. This expedites the removal of accumulated materials from the membrane–solution interfacial region provided by the scouring action of the retentate flowing over the membrane surface.

The combination of the flux and the total membrane area determine the conversion or recovery of the process. The conversion, normally expressed as a percentage Θ, is the amount of the feed that is recovered as permeate. Thus, for a concentration C and flow Q in feed, retentate and permeate (Fig. 2.15), a simple mass balance dictates that:

\[ Q = Q_P + Q_R \]  
\[ QC = Q_P C_P + Q_R C_R \]

where % recovery or conversion is given by:

\[ \Theta = \frac{Q_P}{Q} \]

and the subscripts P and R refer to permeate and retentate, respectively.

![Figure 2.14](image)
Rejection
The permselective property of the membrane is normally quantified as the rejection where:

$$R = 100\% \left(1 - \frac{C_p}{C}\right)$$

(2.4)

It is possible to have negative rejection values if the membrane is selective for specific contaminants, as would be the case for an extractive membrane system.

2.2.2 The driving force

The driving force for the process may be a transmembrane pressure gradient, as with filtration and reverse osmosis (Fig. 2.16), a concentration gradient, as with dialysis, or electromotive, as with electrodialysis. In almost all pressure-driven membrane processes applied to water treatment the desired permeate is water, such that the retained or rejected material (the retentate) is concentrated. In extractive and electrodialytic operations the permeate is the dissolved solute and the retentate the product water. For extractive systems the driving force is a concentration gradient, whereas for electrodialysis an applied potential difference is employed to move dissolved ions through electromigration.

Since the flux and driving force are interrelated, either one can be fixed for design purposes. It is usual to fix the value of the flux and then determine the appropriate value for the transmembrane pressure for pressure-driven processes. An analogous situation arises in the determination of operational parameter values for electrodialysis, where the appropriate operating current density is normally fixed and the voltage determined accordingly. The actual flux or current density value chosen depends upon the desired operating regime.

2.2.3 Factors opposing the driving force

The overall resistance at the membrane–solution interface is increased by a number of factors which each place a constraint on the design and operation of membrane process plant:
the concentration of rejected solute, as in RO and UF, or permeated ions, as in ED, near the membrane surface,

- the depletion of ions near the membrane surface, as with ED,

- the precipitation of sparingly soluble macromolecular species (gel layer formation, as in UF) or salts (scaling, as in RO) at the membrane surface, and

- the accumulation of retained solids on the membrane (cake layer formation, as in MF).

All of the above contribute to membrane fouling. Fouling can take place through a number of physicochemical and biological mechanisms which all relate to increased deposition of solid material onto the membrane surface (also referred to as blinding) and within the membrane structure (pore restriction or pore plugging/occlusion). This is to be distinguished from clogging, which is the filling of the membrane channels with solids due to poor hydrodynamic performance. Fouling may be both temporary (removed by washing) and permanent (removed only by use of chemicals). Since it is intimately related to concentration polarisation, under certain limiting conditions its effects can be determined from a simple theoretical approach (Section 2.3.2). However, fouling by individual components tends to be specific to membrane material and application.

The membrane resistance is fixed, unless its overall permeability is reduced by components in the feed water permanently adsorbing onto or into the membrane. The resistance imparted by the interfacial region is, on the other hand, dependent upon the total amount of fouling material residing in the region. This in turn depends upon both the thickness of the interface, the feedwater composition (and specifically its foulant content) and the flux through the membrane. The feedwater matrix and the process operating conditions thus largely determine process performance.
In the case of a dead-end filtration process, the resistance increases according to the thickness of the cake formed on the membrane, which would be expected to be proportional to the total volume of filtrate passed. For cross-flow processes, this deposition continues until the adhesive forces binding the cake to the membrane are balanced by the scouring forces of the liquid passing over the membrane. All other things being equal, a cross-flow filtration process would be expected to attain steady-state conditions. In practice, only pseudo-steady-state (or stabilised) conditions are attained due to the unavoidable deposition or adsorption of fouling material.

**Concentration polarisation**

Concentration polarisation (CP) is the term used to describe the tendency of the solute to accumulate at the membrane–solution interface within a concentration boundary layer, or liquid film (Fig. 2.17). This layer contains near-stagnant liquid, since at the membrane surface itself the liquid velocity must be zero. This implies that the only mode of transport within this layer is diffusion, which is around two orders of magnitude slower than convective transport in the bulk liquid region. Rejected materials thus build up in the region adjacent to the membrane, increasing their concentration over the bulk value, at a rate that increases exponentially with increasing flux (Section 2.3.2). The thickness of the boundary layer, on the other hand, is determined entirely by the system hydrodynamics, decreasing in thickness when turbulence is promoted.

For pressure-driven processes, the greater the flux, the greater the build-up of solute at the interface: the greater the solute build-up, the higher the concentration gradient: the steeper the concentration gradient, the faster the diffusion. Under normal steady-state operating conditions there is a balance between those forces transporting the water and constituents within it towards, through and away from the membrane. This balance is determined by CP. CP also raises the effective osmotic pressure at the membrane–solution interface, increasing the required transmembrane pressure for operation. It is thus always

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*Figure 2.17 Concentration polarisation*
Membrane technology

37

desirable to suppress CP by promoting turbulence and/or operating at a flux below that at which CP starts to become significant.

CP effects on specific processes are summarised in Table 2.6. All membrane processes are subject to CP, but it is only in specific cases where certain CP phenomena become significant. The importance of elevation of osmotic pressure depends upon the concentration of the rejected solute in the feedwater, since osmotic pressure is directly related to ion concentration by the van't Hoff equation (Section 2.3.9). Thus for nanofiltration processes, where only part of the feedwater ion content is affected by CP, the osmotic pressure elevation effects are commensurately smaller than for RO which rejects ions almost quantitatively. CP also increases the permeation of the rejected materials through the membrane because of the increase in the transmembrane concentration gradient generated. This can affect the permeate water purity, although the effect is minor. The increase in concentration from the bulk solution to the membrane surface can also change the selectivity of the membrane, particularly for nanofiltration, although again the effect is not generally significant.

Scaling presents a more substantial limitation to operation under conditions of CP. Both RO and NF are subject to scaling by divalent salts formed through the CP of ions contributing to hardness. In ultrafiltration, precipitation of sparingly soluble organic solutes produces a gel layer whose permeability and permselectivity often differs from that of the membrane on which it sits. This gel, or dynamic, layer then determines the process performance with respect to both the hydraulics and the product water quality.

In the case of electrodialysis, the effect of polarisation in the liquid film is to deplete ions in that region as ions are extracted through the membrane faster than they arrive at the interface from the bulk solution (Fig. 2.18). Depletion of the permeating ion has two principal effects:

- the electrical resistance increases, and
- the concentration of the permeating ion at the membrane surface of the depletion side decreases to a level that approaches that of the innate water dissociation products (hydroxide and hydrogen ions).

<table>
<thead>
<tr>
<th>Process</th>
<th>Osmotic pressure elevation</th>
<th>Electrical resistance elevation</th>
<th>Scaling</th>
<th>Gel layer formation</th>
<th>Selectivity change</th>
</tr>
</thead>
<tbody>
<tr>
<td>RO</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td>(X)(^a)</td>
</tr>
<tr>
<td>NF</td>
<td>(X)(^a)</td>
<td></td>
<td>X</td>
<td>X</td>
<td>(X)(^a)</td>
</tr>
<tr>
<td>ED</td>
<td>–</td>
<td>X(^2)</td>
<td>(X)(^a)</td>
<td>–</td>
<td>X(^b)</td>
</tr>
<tr>
<td>UF</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>MF</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) More marginal effect.

\(^b\) Depletion polarisation.
Because the membranes used in electrodialysis are generally non-selective for counter-ions, water product ions can pass through the membrane under severe depletion conditions. This phenomenon tends to take place at the anion exchanging membrane because this ion exchange material catalyses the dissociation of water (water splitting) at the low ionic strengths prevailing in the depleted region. The local hydroxide concentration is then effectively increasing and so increasing the passage of hydroxide through the membrane in preference to the contaminant ion (normally chloride). This decreases the electrical efficiency of the process, but more importantly increases the pH on the permeate side of the membrane (the concentrate stream, in this case) which promotes precipitation of hardness salts. CP thus promotes scaling in all dense membrane processes, through the underlying mechanisms differ between pressure-driven and extractive processes. Whilst CP places an upper limit on the flux employed in pressure-driven processes, in the case of electrodialysis its effect is to limit the degree of desalination attainable from a single passage through the stack to below 50%. This is because the rate of desalination is dictated by the current, which is the same at the stack inlet and outlet. It follows that if a sufficient current is applied to remove 50% of the ions at the inlet of the stack, the limiting condition of zero concentration will be reached at the stack outlet on the diluate side.

The relationship between driving force and polarisation in pressure-driven membrane separation processes can be summarised as follows:

- The flow through a given type of membrane varies as the membrane area and the net applied driving force; and the power consumption is proportional to the driving force, and inversely proportional to the membrane area installed. This is analogous to electrical conduction, where the current varies with the cross-sectional area of copper in the cable and with the applied voltage, and the power loss in the cable varies with the voltage loss and inversely with the area.
- The selective nature of the process means that rejected material remains on the membrane surface. Cross-flow operation affords some limitation to
the extent to which rejected material accumulates in the interfacial region.

These two factors are, of course, interlinked: a high driving force yields high flux and a high rate of rejected material collecting on the membrane surface, which then needs to be dispersed rapidly if the process is not to grind to a halt. In extractive and dialytic processes, CP tends to deplete the permeating species at the membrane, which in electrodialysis has the effect of increasing electrical resistance and decreasing permselectivity.

### 2.2.4 Critical flux

The critical flux concept was originally presented by Field et al. (1995). These authors stated that: “The critical flux hypothesis for microfiltration is that on start-up there exists a flux below which a decline of flux with time does not occur; above it, fouling is observed”. Two distinct forms of the concept have been defined. In the strong form, the flux obtained during sub-critical flux is equated to the clean water flux obtained under the same conditions. However, clean water fluxes are rarely attained for most real feedwaters due to irreversible adsorption of some solutes. In the alternative weak form, the sub-critical flux is the flux rapidly established and maintained during the start-up of the filtration, but does not necessarily equate to the clean water flux. Alternatively, stable filtration operation, i.e. stable permeability for an extended period of time, has been defined as sub-critical operation even when preceded by an initial decline in flux (Howell, 1995). Such conditions would be expected to lead to lower critical flux values than those obtained for absolutely constant permeability operation (i.e. from \( t=0 \)), however, since an initial permeability decline implies foulant deposition.

A number of slightly different manifestations of sub-critical flux operation have been proposed, largely depending on the method employed. The most microscopically precise definition equates the critical flux to that flux below which no deposition of colloidal matter takes place. Kwon and Vigneswaran (1998) equated the critical flux to the lift velocity as defined by lateral migration theory (Table 2.12: Section 2.3.2), as introduced by Green and Belfort (1980). This rigorous definition is difficult to apply because of the relative complexity of the determination of the lift velocity, particularly for heterogeneous matrices. On the other hand, experimental determination of critical flux by direct observation of material deposition onto the membrane has been conducted using model homodispersed suspensions of polystyrene latex particles (Kwon and Vigneswaran, 1998), and some authors have also used mass balance determinations (Kwon et al., 2000).

Given the limitations of applying particle hydrodynamics to the identification of the critical flux in real systems, recourse generally has to be made to experimental determination. By plotting flux against the transmembrane pressure it is possible to observe the transition between the linearly pressure-dependent flux and the onset of fouling, where deviation from linearity
commences. The flux at this transition has been termed "secondary critical flux" (Bouhabila et al., 1998). However, whilst potentially useful in providing a guide value for the appropriate operating flux, the absolute value of the critical flux obtained by this method is likely to be dependent on the exact method employed and, specifically, the rate at which the flux is varied with time. A common practice is to incrementally increase the flux for a fixed duration for each increment, giving a stable TMP at low flux but an ever-increasing rate of TMP increase at higher fluxes. This flux-step method defines the highest flux for which TMP remains stable as the critical flux. This method is preferred over the TMP-step method since the former provides a better control of the flow of material deposition on the membrane surface, as the convective flow of solute towards the membrane is constant during the run (Defrance and Jaffrin, 1999a). However, no single protocol has been agreed for critical flux measurement, making comparison of reported data difficult. It is also becoming apparent that irreversible fouling can take place at operation below the critical flux (Cho and Fane, 2002).

2.3 The theory

There are essentially two approaches to describing mass transport in membrane processes. The simplest is to add the hydraulic resistance of the membrane to that of the cake or fouling layer to determine the relationship between the flux and pressure through simple empirical Darcian relationships. This approach relies on a knowledge of the resistance of both membrane and fouling/cake layer. The membrane resistance can be determined either directly from ex situ experimental measurements using pure water or, in the case of porous membranes, through well-understood fluid physics relationships. Provided the cake or fouling layer can be measured empirically, and assumptions can be made about the effect of operation on the bulk membrane permeability, resistance theory can be usefully applied to determine hydraulic relationships without recourse to further theoretical development. It is, indeed, common practice to refer to the membrane and cake or fouling layer resistance when defining filtration operation. Other cake filtration empirical models are based on deposition of solids within the membrane pores, thus accounting for the change in the permeability with time but more as a diagnostic tool than for predictive purposes.

The development of predictive models for membrane mass transfer from first principles is much more problematic and relies on mathematical description of the system hydrodynamics, membrane structure and feedwater matrix. It is only under specific limiting conditions that simple analytical expressions can be used with complete impunity (for example dense, homogeneous, i.e. non-porous membranes where the water content of the membrane is small, or isoporous, non-adsorptive uncharged membranes). Hence, the combination of solution–diffusion and sieving mechanisms (or more specifically capillary flow), as exists in nanofiltration processes, substantially complicates the mathematical description. Moreover, no simple fundamental analytical expressions can account for fouling, in particular internal or permanent fouling where adhesive
forces become important. Instead, simple experimental measurements are used to assess the fouling propensity of feedwaters for specific membrane processes, and dense membrane processes in particular (Section 2.4.3).

Notwithstanding the difficulties involved, a number of models have been presented to define the operational determinants of various membrane processes. To describe the derivation of each of these in detail would be beyond the scope of this review since the outcomes, i.e. the ultimate analytical expressions generated, are often specific to the process under consideration and the assumptions made. The starting point for the theoretical development of water and solute flows through a dense membrane varies according to the relative importance placed of solution–diffusion, sorption, pore flow and electrical charge. Most of the mathematical derivations of system hydrodynamics for cross-flow operation are based on film theory (which incorporates the concentration polarisation model). Film theory assumes the interfacial region mass transport to be determined by the degree of concentration polarisation, which can then be calculated from the fluid cross-flow, the permeate flux and the solute diffusivity.

2.3.1 Membrane mass transfer control

Under the simplest operational conditions, the resistance to flow is offered entirely by the membrane. For porous membrane systems, the flux can be expressed as:

\[ J = \frac{\Delta p}{\mu R_m} \]  \hspace{1cm} (2.5)

where \( J \) is the flux in m s\(^{-1}\), \( \Delta p \) is the transmembrane pressure, \( \mu \) is the fluid viscosity and \( R_m \) is the resistance of the membrane in m kg\(^{-1}\). For microporous membranes, specifically those used for microfiltration, the Hagen–Poiseuille equation may be considered applicable for a permeate undergoing laminar flow through cylindrical pores. The resistance \( R_m \) then equates to:

\[ R_m = \frac{K(1 - \varepsilon_m)^2 S_m^2 l_m}{\varepsilon_m^3} \]  \hspace{1cm} (2.6)

where \( \varepsilon \) is the porosity (or voidage), \( S_m \) the pore surface area to volume ratio and \( l_m \) the membrane thickness. \( K \) is a constant equal to 2 for perfectly cylindrical pores but changes for other geometries. It is apparent from Equation (2.5) that temperature has a profound impact upon the flux through the viscosity, which increases by around 3% for each degree drop in temperature below 25°C.

For dense membranes a number of expressions have been developed (Table 2.7) which are derived from a variety of approaches:

- solution–diffusion,
- sorption–capillary flow,
- Donnan equilibrium,
Table 2.7 Dense membrane rejection expressions based on different models (based on Bhattacharyya and Williams, 1992)

<table>
<thead>
<tr>
<th>Approach</th>
<th>Rejection</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solution–diffusion</td>
<td>( \left( 1 + \frac{K_s}{K_w} \left( \frac{1}{\Delta P - \Delta \pi} \right) \right) ) (^{-1} )</td>
<td>Lonsdale et al. (1965)</td>
</tr>
<tr>
<td>Solution–diffusion imperfection</td>
<td>( \left( 1 + \frac{K_s}{K_w} \left( \frac{1}{\Delta P - \Delta \pi} \right) \right) ) (^{-1} ) ( \left( 1 + \frac{K_s}{K_w} \left( \frac{1}{\Delta P - \Delta \pi} \right) \right) ) (^{-1} )</td>
<td>Sherwood et al. (1967)</td>
</tr>
<tr>
<td>Surface force pore flow</td>
<td>( 1 - \frac{\int_0^1 C_p v_p r_p d r_p}{\int_0^1 v_p r_p d r_p} )</td>
<td>Matsuura and Sourirajan (1981); Sourirajan and Matsuura (1985)</td>
</tr>
<tr>
<td>Donnan equilibrium</td>
<td>( 1 - \left( \frac{C_p}{Q_m} \right)^y \left( \frac{\gamma}{\gamma_m} \right)^{y-1} ) (^{1/2} )</td>
<td>Bhattacharyya and Chen (1986)</td>
</tr>
</tbody>
</table>

\( ^{a} \) b, dimensionless frictional force (frictional force acting on solute in pore vs. that acting on solute in bulk solution); \( C_w \) concentration of water in the membrane; \( C_s \), \( \exp(y) / (1 + \exp(y - 1)(b_v / \exp(z))) / \Delta C_s \), solute concentration difference across the membrane = \( C_{s1} - C_{s2} \); \( C_{s1,s2} \), solute concentration on feed, permeate side of membrane; \( C_p \), concentration of co-ion in bulk solution; \( D_w \), (Brownian) diffusion coefficient of water in the membrane; \( D_s \), (Brownian) diffusion coefficient of solute in the membrane; \( K_w \), water permeability in the membrane; \( D_p C_w V_w / RT \); \( K_s \), solute permeability in the membrane; \( D_p \kappa_s; K_p \), empirical coefficient; \( Q_m \), charge capacity of membrane; \( R \), gas constant; \( r_p \), dimensionless radial distance (\( r / r_p \)); \( T \), absolute temperature; \( v \), dimensionless radial velocity (\( v_r = v / (K_v - 1) \)); \( V_w \), partial molar volume of water; \( y \), \( z \), charge on species \( y \) (the co-ion) and \( z \) (counter-ion) respectively; \( Y_p / Z_i \), \( (X^{z+} + Z^+) \); \( Y_m \), activity coefficients in solution and membrane respectively; \( \kappa_m \) solute distribution coefficient; \( p \), osmotic pressure; \( \Phi_e \), dimensionless electrostatic or van der Waals forces parameter.

- extended Nernst–Planck equation, and
- irreversible thermodynamics.

Pure solution–diffusion is based upon the assumption that both the solute and solvent (i.e. water) dissolve in a homogenous non-porous membrane and then diffuse across it as discrete (uncoupled) materials. The overall rate of transfer (i.e. flux) of each, and thus the purity of the permeate product, depends upon their relative solubilities and diffusivities (Lonsdale et al., 1965):

\[
\text{Solvent (water) flux:} \quad \frac{K_w}{\lambda} (\Delta P - \Delta \pi) \tag{2.7}
\]

\[
\text{Solute (salt) flux:} \quad \frac{K_s}{\lambda} (\Delta C_s) \tag{2.8}
\]

where \( \Delta P \) and \( \Delta \pi \), respectively, represent the applied transmembrane and osmotic pressures. \( \Delta C_s \) is the solute concentration difference across the membrane. \( K_w \) and \( K_s \) represent the water and solute permeabilities in the membrane and \( \lambda \) is the membrane thickness. The ratios \( K_w / \lambda \) and \( K_s / \lambda \) are
normally denoted \( A \) and \( B \), respectively. According to this approach, the transport of solute is not directly related to transmembrane pressure.

The osmotic pressure \( \Pi \) is given by the well-known van't Hoff equation:

\[
\Pi = yRT \sum C_i
\]  

where \( R \) is the gas constant, \( T \) the absolute temperature, \( \Sigma C_i \) the total concentration of individual solute ions and \( y \) the osmotic coefficient accounting for non-ideal behaviour. The coefficient \( y \) normally takes a value between 0.7 in concentrated solutions and unity at infinite dilution, and equates to the mean of the activity coefficients of the dissolved ions and the degree of dissociation. However, the relationship between \( \Pi \) and \( C \) is rarely linear, as Equation (2.9) suggests, due to departure from ideality at high concentrations.

The simple solution–diffusion model can be extended to account for pore flow by adding an extra empirical coefficient to the water and solute flux expressions. Use of the model is, however, constrained by the necessity of determining this coefficient experimentally. The model can also be extended to allow coupling of solute and solvent flows, although this also introduces empirical constants.

Pore flow models assume that the membrane is microporous with cylindrical pores. The membrane material is assumed to preferentially adsorb water, as a monolayer, both within the pores and at the surface. The water is then transported across the membrane at a rate depending upon chemical interaction, frictional and diffusive forces. The expression can be modified to account for steric and electrostatic effects, and can take account of the pore size distribution of the membrane. Charged pores tend to increase the effective viscosity of the water in the pores due to electroviscous effects, but these effects alone do not appear to account for reported charged membrane permeability changes with ionic strength and pH.

The Donnan equilibrium model takes account of solute charge and membrane charge which, through repulsion of co-ions and attraction of counter-ions, creates a Donnan potential at the membrane surface. This has the effect of repelling co-ions and, since electroneutrality must be preserved, counter-ions are also rejected. This approach applies to both electrodialysis and nanofiltration and can provide qualitative trends, but does not allow a quantitative description of nanofiltration since it does not take account of diffusive and convective transport. The extended Nernst–Planck equation has also been used to describe transport in charged membranes, but with similar limitations.

### 2.3.2 Fouling/cake layer mass transfer control

**Resistance model: cake filtration**

The simplest way of accounting for the additional resistance offered by the material accumulating in the interfacial region is to simply add its resistance \( R_c \) (the cake layer resistance) to that of the membrane, such that Equation (2.5) becomes:
where, in the case of dead-end operation, \( R_c \) is time dependent. However, this simple equation is only useful if there is prior knowledge concerning the hydraulics of the cake or fouling layer. In the model case of the development of an incompressible cake of homodispersed granular material the cake resistance follows the same form as Equation (2.6):

\[
R_c = \frac{K'(1 - \varepsilon_c)^2 S_c^2}{\varepsilon_c^3}
\]  

(2.11)

where the symbols refer to the same parameters as before with reference to the filter cake. In this case, however, \( K' \) takes a value of 5 for spherical (or neo-spherical) geometry (Grace, 1956), and the equation – or more usually the pressure gradient form derived from it – represents an expression of the well-known Kozeny–Carman equation.

Whilst the above equations may be used to calculate the total resistance to filtration, their use is constrained by a number of simplifying assumptions:

- flow is considered only in the direction orthogonal to the membrane: tangential movement is ignored,
- suspended particles are assumed homodispersed (i.e. all the same size, shape) and neutrally buoyant (i.e. non-sedimenting),
- particles are considered incompressible,
- the filter cake is also considered incompressible,
- cake porosity, hence permeability, is assumed to be independent of time, and
- migration of particles through the cake with time is ignored.

In fact, filtration can be characterised on the basis of more than just cake filtration behaviour. Four filtration models, originally developed for dead-end filtration (Grace, 1956), have been proposed to describe the initial flux decline. All models imply a dependence of flux decline on the ratio of the particle size to the pore diameter (Table 2.8). The standard blocking and cake filtration models appear most suited to predicting initial flux decline during colloid filtration (Visvanathan and Ben Aim, 1989) or protein filtration (Bowen et al., 1995). According to Bowen and co-workers, four consecutive steps had been defined: (1) blockage of the smallest pores, (2) coverage of the larger pores inner surface, (3) superimposition of particles and direct blockage of larger pore and (4) creation of the cake layer. All of the models contain empirically derived parameters (\( A \) and \( B \) in Table 2.8), although some have been refined to incorporate other key determinants (Table 2.9). On the other hand, a number of empirical and largely heuristic expressions have been proposed for particular matrices and/or applications, for example the filtration of activated sludge by membrane bioreactors (Table 2.10). Classical dead-end filtration models can be
adapted for cross-flow operation if the proportion of undeposited solute material can be calculated.

The expressions listed in Table 2.10 illustrate a common difficulty in describing filtration behaviour. Even for a supposedly generic matrix (mixed liquor), the innate heterogeneity and diversity in composition of real feedwater matrices forbids the formulation of general equations describing filtration behaviour, even under steady-state conditions. Different key water quality determinants (chemical oxygen demand, dissolved organic carbon and mixed liquor (volatile) suspended solids) have been identified as the basis for quantifying fouling propensity and dynamic behaviour. Most recent publications of experimental studies of MBR processes make little or no reference to dynamic modelling, with data interpretation being substantially limited to reporting of hydraulic resistance values (Choo and Lee, 1996; Chang and Lee, 1998; Defrance and Jaffrin, 1999b). The more global semi-empirical models that have been developed (Nagaoka et al., 1998), as with many other semi-empirical models in this area, rely on specific hydraulic resistance data which is likely to be specific to the system under investigation.

Dead-end filtration theory is thus to a very large extent limited by the accuracy of the representation of the interaction between the solids and the membrane material. The same also applies to cross-flow filtration. However, for dense membrane processes where the water can be considered to be a continuum with no complicating solid-liquid interactions, modelling from first principles of the resistance offered by the membrane-solution interface, due to concentration polarisation, is possible.

**Classical concentration polarisation model**

As already stated (Section 2.2.3), concentration polarisation describes the tendency of the solute to build up in the membrane solution interfacial region, and the extent to which this occurs depends on:

<table>
<thead>
<tr>
<th>Approach</th>
<th>Physical cause</th>
<th>Description</th>
<th>Equationa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cake filtration</td>
<td>Boundary layer resistance</td>
<td>Deposit of particles larger than the membrane pore size onto the membrane surface</td>
<td>(\frac{t}{V} = AV - B)</td>
</tr>
<tr>
<td>Complete blocking</td>
<td>Pore blocking</td>
<td>Occlusion of pores by particles with no particle superimposition</td>
<td>(-\ln(J/J_0) = At + B)</td>
</tr>
<tr>
<td>Intermediate</td>
<td>Long-term adsorption</td>
<td>Occlusion of pores by particles with particle superimposition</td>
<td>(\frac{1}{f} = At + B)</td>
</tr>
<tr>
<td>Standard blocking</td>
<td>Direct adsorption</td>
<td>Deposit of particles smaller than the membrane pore size onto the pore walls, reducing the pore size</td>
<td>(\frac{t}{V} = At + B)</td>
</tr>
</tbody>
</table>

\(A, B, \) constants (value dependent upon cake and system characteristics); \(V\). cumulative volume of permeate at time \(t\); \(J_0, J\). flux initially and at time \(t\) respectively.
### Table 2.9 Expressions developed to describe, for dynamic behaviour, dead-end and cross-flow filtration

<table>
<thead>
<tr>
<th>Model</th>
<th>Equationa</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dead-end filtration</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pore blocking model</td>
<td>[ J = J_0 \exp\left(\frac{-a_{p0}A}{n_p}t\right) ]</td>
<td>Hermia (1982)</td>
</tr>
<tr>
<td>Pore constriction model</td>
<td>[ J = \frac{J_0}{\left(1 + \frac{a_{p0}J_0}{n_p\pi\delta_m r_p^2}t\right)^\frac{1}{2}} ]</td>
<td>Hermia (1982)</td>
</tr>
<tr>
<td>Cake formation model</td>
<td>[ J = \left(\frac{1 + 2\delta_d R_e^2 C_0 A \Delta P}{v_a R_m^2 t}\right)^{1/2} ]</td>
<td>Hermia (1982)</td>
</tr>
<tr>
<td><strong>Cross-flow filtration</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shear-induced diffusion from thick layers</td>
<td>[ J = J_0 \left(1 - \frac{2J_0 R_e C}{R_m (C_e - C)} t\right)^{-1/2} ]</td>
<td>Romero and Davis (1988)</td>
</tr>
<tr>
<td>Brownian diffusion from thick layers</td>
<td>[ \langle J \rangle = 1.31 \left(\frac{D^2 \gamma_0}{L}\right)^{1/3} \left(\frac{C_e}{C} - 1\right)^{1/3} ] for (t &lt; t_{ss})</td>
<td>Song (1998), after Song and Elimelech (1995)</td>
</tr>
<tr>
<td></td>
<td>[ \langle J \rangle = \frac{1}{L} \int_0^X [J_s(x)dx - {L - X(t)}J(t)] ] for (t &lt; t_{ss})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[ X(t) = 4.81(D^2 \gamma_0) \left(\frac{C_e}{C} - 1\right) \left(\frac{C}{C_e}\right)^{1/2} \left(\frac{R_e^2 t}{\Delta P - \Delta P_e}\right)^{-3/2} ]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[ \Delta P_e = \frac{3kT}{4\pi R^3 N_p} ]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[ J(t) = \left(\frac{\Delta P - \Delta P_e}{R_m}\right)^{1/2} \left(1 + \frac{2R_e^2 \Delta P - \Delta P_e}{R_m^2 C_e} \right)^{1/2} ]</td>
<td></td>
</tr>
</tbody>
</table>

*a* \(a_{p0}\), no. of pores blocked per filtrate volume passed; \(a_{p0p}\), foulant volume deposited within pores per filtrate volume passed; \(a_{depp}\), fraction of foulant depositing on membrane; \(A\), filter area; \(J_0\), \(J_s\), flux initially, at time \(t\), at equilibrium; \(n_p\), no. of pores; \(r_p\), pore radius; \(R_m\), membrane resistance; \(P\), operating pressure difference; \(P_e\), critical pressure difference (pressure required for cake to form); \(t\), filtration time; \(t_{ss}\), steady-state filtration time (time after which flux is constant); \(X\), distance of the front of the dynamic layer from the entrance; \(n_p\), bulk suspension viscosity.

- the propensity of the rejected (solute) species to diffuse (i.e. their diffusivity),
- the notional thickness of the stagnant region, and
- the rate at which solute species are added to the stagnant region.

The first of these relates to the solute itself, and principally to its size. The latter two factors, on the other hand, are determined mainly by operating conditions. The thickness of the stagnant region \(\delta\) can be determined from:
<table>
<thead>
<tr>
<th>Application/derivation</th>
<th>Expression*</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein</td>
<td>[ J = \frac{\Delta P}{\eta_0 [R_m - R'_b M_c (1 - e^{-k_1 t}) + R'_b M_b]} ]</td>
<td>Fane (1986)</td>
</tr>
<tr>
<td></td>
<td>[ J = \frac{\Delta P}{\eta_0 (k_2 U - 2.5 + k_1) R'_b U} ]</td>
<td>Fane (1986)</td>
</tr>
<tr>
<td>Classical cake filtration</td>
<td>[ J = \frac{\Delta P}{\mu (R_m + \alpha C_{MLSS})} ]</td>
<td>Chang <em>et al.</em> (2001); Shimizu <em>et al.</em> (1993)</td>
</tr>
<tr>
<td>Concentration polarization</td>
<td>[ J = a \cdot b \log C_{DOC} ]</td>
<td>Ishiguro <em>et al.</em> (1994)</td>
</tr>
<tr>
<td>Sidestream MBR</td>
<td>[ J = \frac{\Delta P}{\mu (R_m + 843 \Delta P_{COD} C_{MLSS} C_{MLVSS} (\delta \log 10 R'_b \cdot k_1 \cdot d_k))} ]</td>
<td>Sato and Ishii (1991)</td>
</tr>
<tr>
<td>Sidestream MBR</td>
<td>[ J = J_0 \exp \left( \frac{k \Re (C_{MLSS} - C_{MLVSS})}{C_{MLVSS}} \right) ]</td>
<td>Krauth and Staab (1993)</td>
</tr>
<tr>
<td>HF submerged MBR</td>
<td>[ J = k U_a \Phi C_{MLSS} ]</td>
<td>Shimizu <em>et al.</em> (1996)</td>
</tr>
</tbody>
</table>

* a, b, k, empirical constants; C, concentration; \( J_0 \), initial flux; \( k_{1,2,3} \), empirical first-order rate constants; \( M_b \), boundary layer cake deposit; \( M_c^* \), maximum cake deposit (mass per unit area); \( R'_b \), specific boundary layer resistance; \( U_a \), approach velocity, air; \( \varphi \), membrane geometric hindrance factor (membrane geometry dependent). Subscripts: COD, chemical oxygen demand; DOC, dissolved organic carbon; ML(V)SS, mixed liquor (volatile) suspended solids.

- the physical properties of the liquid, which for most water treatment processes change only marginally with chemical water quality and can normally be accurately expressed as a function of temperature,
- the shape and size of the flow channels within the module, and
- the mean velocity of liquid flowing through the channels.

The rate at which ions accumulate in the stagnant film is simply determined by the flux and the rejection. Therefore, provided the system is well defined, the degree of concentration polarization can be calculated and its effects on the operation of the membrane process assessed. Mathematical description proceeds by conducting a material balance at the membrane, where the build-up of solute at the interface is countered by the diffusive flux of solute away from membrane. This essentially entails the balancing of four fluxes:

- back diffusion of solute away from the membrane by Fick's first law of diffusion, which states that the diffusion rate is proportional to the concentration gradient,
- convective transport of the solute to the membrane,
- convective diffusion of the solute through the membrane, and
- convective diffusion of permeate through membrane.

Assuming a one-dimensional system (i.e., no longitudinal mass transfer) and a constant value \( \delta \) for the boundary layer thickness (Fig. 2.18), the concentration polarization under steady-state conditions based on film theory can be defined as:
\[ J = \frac{D}{\delta} \ln \left( \frac{C^*}{C} \right) \]  

(2.12)

where \( D \) is the diffusion coefficient in m\(^2\) s\(^{-1}\) and \( C^* \) and \( C \) are the respective concentrations at the membrane surface and in the bulk solution. The ratio \( C^*/C \) is a critical term, sometimes denoted \( \varphi \) or \( \beta \), and referred to as the concentration polarisation parameter or index. Note that Equation (2.12) includes no pressure term, although the transmembrane pressure is inferred by the flux value. In the case of gel layer formation, the term \( C^* \) equates to the concentration of solute in the gel layer, for which some experimentally determined values have been reported for specific components (Cheryan, 1998).

Determination of the flux from Equation (2.12) relies on knowledge of the solute diffusivity, the boundary layer thickness and the solute concentration at the membrane surface. If the solute comprises dissolved ions or small molecules, as would be the case for pressure-driven dense membrane processes, then \( D \) is simply given by the Stokes–Einstein equation:

\[ D = \frac{2k_B T}{3\pi \mu r_p} \]  

(2.13)

where \( k_B \) is the Boltzmann constant, \( T \) is absolute temperature and \( r_p \) is the solute radius.

Con conventionally, the ratio \( D/\delta \) is assigned the mass transfer coefficient \( k \) and the various operational and performance determinants expressed in terms of the non-dimensionalised groups of \( Sh \) (Sherwood number), \( Re \) (Reynolds number) and \( Sc \) (Schmidt number):

\[ J = k \ln \left( \frac{C^*}{C} \right) \]  

(2.14)

\[ k = ShD/d \]  

(2.15)

\[ Sh = aRe^bSc(d/L)^n \]  

(2.16)

\[ Re = \rho Ud/\mu \]  

(2.17)

\[ Sc = \mu/\rho D \]  

(2.18)

where \( \rho \) is the fluid density, \( U \) its velocity, \( d \) and \( L \) the hydraulic dimension and length of the membrane and \( a, b, c \) and \( n \) are constants. The hydraulic dimension \( d \) is equal to four times the ratio of the channel cross-section to its perimeter, and is thus equal to the diameter of a tube or twice the height of a wide parallel flow channel.

Appropriate values for \( a, b, c \) and \( n \), based on Newtonian behaviour, are given in Table 2.11. The solution for laminar flow and Brownian diffusive transport is attributed to Lévêque (Lévêque, 1928; Porter, 1972). The Lévêque solution assumes channel flow with completely impermeable boundaries, and is thus strictly only applicable to membrane permeation systems with a permeate flux.
much smaller than the cross-flow velocity, which is normally the case in cross-flow membrane systems. However, all membrane elements include a spacer that provides a degree of turbulence promotion, which can be accounted for through the inclusion of additional terms in Equation (2.16) containing parameters characterising the spacer. An example, which incorporates a term in the spacer mesh width parameter $m$ (Chiolle et al., 1978), is included in Table 2.11.

Expressions can also be presented in terms of the shear rate $\gamma$ (Table 2.12). The shear rate is given by the ratio of the cross-flow velocity $U$ to the hydraulic dimension and is geometry dependent. For parallel flow channels of height $h$ its maximum value is given by:

$$\gamma_0 = 6U/h$$  \hspace{1cm} (2.19)

For tubes of diameter $d$ the equivalent expression is:

$$\gamma_0 = 8U/d$$  \hspace{1cm} (2.20)

The impact of concentration polarisation on process operation depends upon the process. For pressure-driven processes the main impacts are on feedwater osmotic pressure and scalant levels, since both will increase by a factor of up to $\varphi/(1 - \theta)$ overall. The osmotic pressure effect raises the required applied pressure to maintain the target flux in accordance with Equation (2.7), whereas the increase in scalant levels may produce precipitation of these salts. This demands an appropriate choice of process operating conditions (Section 2.4.2).

In the case of extractive processes, and electrodialysis specifically, polarisation depletes the ions in the boundary layer and places an upper limit on the operating current density, and so on the operating current (Section 2.4.2). The current density attained under this condition is known as the limiting current density $I_{\text{lim}}$:

$$I_{\text{lim}} \sim 2kCF$$  \hspace{1cm} (2.21)

where $k$ is the mass transfer coefficient given by Equations (2.15)–(2.18). $C$ is the bulk equivalent concentration (in eq m$^{-3}$) of the diluate (i.e. the desalinated product) and $F$ is the Faraday constant, which takes a value of 96 500 C eq$^{-1}$.

Table 2.11  Examples of empirical and semi-empirical values of constants in Equation (2.16)

<table>
<thead>
<tr>
<th>Flow</th>
<th>Characteristic constants</th>
<th>Additional term</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$a$</td>
<td>$b$</td>
</tr>
<tr>
<td><strong>Open channel</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Turbulent</td>
<td>0.023 or 0.065</td>
<td>0.875</td>
</tr>
<tr>
<td></td>
<td>0.04</td>
<td>0.75</td>
</tr>
<tr>
<td>Laminar (Lévêque)</td>
<td>1.62 or 1.86</td>
<td>0.33</td>
</tr>
<tr>
<td><strong>Filled channel</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intermediate flow</td>
<td>1.065</td>
<td>0.5</td>
</tr>
<tr>
<td>Model</td>
<td>Equation</td>
<td>Reference</td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>-----------</td>
</tr>
<tr>
<td>Lévéque solution: laminar flow. Brownian diffusive transport, ( J \perp U )</td>
<td>( J = 0.807 \left( \frac{D_B \gamma_0}{L} \right)^{1/3} \ln \left( \frac{C^*}{C} \right) )</td>
<td>Porter (1972), after Lévéque (1928)</td>
</tr>
<tr>
<td>Similarity solution for laminar flow. Brownian diffusive transport, ( J \rightarrow U )</td>
<td>( J = 1.31 \left( \frac{D_B \gamma_0 C^*}{CL} \right)^{1/3} )</td>
<td>Davis and Sherwood (1990) Romero and Davis (1988)</td>
</tr>
<tr>
<td>Fully developed turbulent flow</td>
<td>( J = 0.024 \left( \frac{\gamma_0^{1.4} D_B^{0.67}}{\rho^{0.2} \mu^{0.47}} \right)^{1/3} \ln \left( \frac{C^*}{C} \right) )</td>
<td>Porter (1972)</td>
</tr>
<tr>
<td>Lévéque solution for shear-induced diffusion (based on ( D_S = 0.03 r^2 \gamma_0 ))</td>
<td>( J = 0.078 \left( \frac{C^<em>}{L} \right)^{1/3} \gamma_0 \ln \left( \frac{C^</em>}{C} \right) )</td>
<td>Zydney and Colton (1986), after Ecstein et al. (1977)</td>
</tr>
<tr>
<td>Similarity solution for shear-induced diffusion (based on ( C^* \approx 0.6 ) by volume and ( C &lt; 0.1 ) by volume)</td>
<td>( J = 0.072 \gamma_0 \left( \frac{C^*}{C_L} \right)^{1/3} )</td>
<td>Davis and Sherwood (1990)</td>
</tr>
<tr>
<td>Integral model for shear-induced diffusion from thick layers (based on ( D_S(C) ))</td>
<td>( J = 1.31 \gamma_0 \delta(C) \left( \frac{r Q_{cr}(C)}{LC} \right)^{1/3} )</td>
<td>Romero and Davis (1988)</td>
</tr>
<tr>
<td>Inertial lift velocity (based on thin layers, such that ( J = v_L ))</td>
<td>( J = \frac{0.036 \gamma^2}{16 \rho} )</td>
<td>Drew et al. (1991)</td>
</tr>
<tr>
<td>Surface transport</td>
<td>( J = 2.4 r \gamma_0 (r^2 R_c')^{3/5} \cot \theta )</td>
<td>Song and Elimelech (1995)</td>
</tr>
</tbody>
</table>

\( D_B, D_S \) diffusion coefficient for Brownian and shear-induced diffusion respectively; \( \gamma_0 \), maximum shear rate; \( C^*, C \) concentration at the membrane-solution interface and the bulk retentate solution respectively; \( C \), dimensionless concentration; \( L \), membrane elemental length; \( U \), cross-flow velocity; \( \nu \), kinematic viscosity (\( \eta/\rho \)); \( r \), particle radius; \( \eta(C) \), concentration-dependent dimensionless viscosity; \( Q_{cr}(C) \), concentration-dependent excess particle flux; \( R_c' \), specific cake resistance (\( R_0/cake thickness \)); \( v_L \), inertial lift velocity; \( \cot \theta \), surface morphology parameter.

**Modified concentration polarisation model**

Complications arise in the concentration polarisation model when it is applied to systems in which colloidal and/or suspended material is present and accumulates in the hydrodynamic boundary layer. In such cases Newtonian behaviour cannot be assumed within or at the outer boundary of the stagnant film, and the deviation from classic film theory increases with increasing solute concentration in the boundary layer. This implies that a correction for non-Newtonian behaviour is needed to account for the local solute concentration-dependent changes in:
• viscosity of the fluid,
• diffusivity of the solute, and
• permeability of the cake.

Some of the analytical expressions developed for the equilibrium length-averaged flux are included in Table 2.12. The solution varies according to the approach taken and the assumptions made, but the general trend is for transport of solute away from the membrane to be much higher than that predicted by Brownian diffusion. This changes the degree of dependency of flux on both cross-flow velocity (and so shear rate, according to Equations (2.18) and (2.19)) and particle size changes significantly from the Brownian diffusion Lévéque model if either shear-induced diffusion or inertial lift are included. For example, flux dependency on cross-flow changes from $U^{0.33}$ proportionality to direct proportionality for shear-induced diffusion (Zydney and Colton, 1986) or $U^2$ for inertial lift (Drew et al., 1991). Kim and Park (1999) based their prediction of critical flux conditions on shear-induced diffusion.

A comprehensive, and complex, solution for equilibrium flux in CFMF has been presented by Romero and Davis (1988). This model accounts the effects of shear-induced diffusion on a non-uniform filter cake whose thickness increases with axial membrane (channel) distance. This is considerably more complex than the model for thin cake deposits (Zydney and Colton, 1986), where a uniform cake deposit over the whole membrane area is envisaged, and requires that the solute concentration dependency of viscosity and diffusivity be predetermined. More recently, the Romero and Davis model has been slightly simplified by basing the cake layer resistance on the Kozeny–Carman equation (Ould-Dris et al., 2000).

Practical verification of modified models
Experimental studies on model colloidal or particulate systems have shown close agreement between experimentally measured steady-state flux data and those predicted from shear-induced diffusive mass transport theory for ideal systems, such as latex, blood, bacterial and fractionated clay suspensions (Zydney and Colton, 1986). The model of Kim and Park (1999) for predicting critical flux, again based on shear-induced diffusion, appears to corroborate well with experimental data from calcium carbonate filtration, the critical flux increasing linearly with particle size. Inertial lift, on the other hand, would appear to be restricted in importance to high shear rates and/or large particles (Davis, 1992).

The shear-induced diffusion model for thick cake layers (Romero and Davis, 1988, 1990) was found to give a reasonable representation of both dynamic and steady-state behaviour for both rectangular channels and ceramic tubular membranes challenged with homodispersed spheres of 0.45–1.37 μm (Romero and Davis, 1991). Ould-Dris et al. (2000), using a slightly simplified adaptation of the Romero and Davis model, also found reasonable agreement between theoretical and experimental steady-state flux for their trials on the less idealised system of granular calcium carbonate. Agreement was only obtained, however,
after allowance was made for differing mean particle size of the cake layer and the bulk suspension, which required experimental verification. A similar limitation appears to apply to the Song model (Song and Elimelech, 1995): an excellent fit with theoretically predicted flux was obtained for CFMF of 0.06 μm homodispersed colloidal silica once the specific cake resistance $R'_c$ and the cake concentration $C_c$ had been surmised (Wang and Song, 1999).

It is generally the case that the modified concentration polarisation models developed to define dynamic and steady-state behaviour during microfiltration are very sensitive to key parameters pertaining to particle transport and the hydraulic resistance of the cake. These parameters are either only calculable for highly idealised systems or else must be determined empirically. For real feedwaters, which may be physically, chemically and biologically complex, and the useful employment of these models for predictive purposes is probably restricted to defining trends. Even then, there are pronounced differences in reported data. For example for a single matrix of bovine blood, flux dependency on cross-flow appears to vary from $U^{0.47}$ to direct proportionality (Cheryan, 1998). Moreover, there appears to be a paucity of mechanistic information on fouling and dynamic behaviour of key heterogeneous matrices, such as natural organic matter (NOM) in surface waters or biomass in membrane bioreactor liquors, that could be used to form the basis of a predictive model.

2.4 Process design and operation

Key elements of process design that contribute to costs are:

- staging,
- power consumption,
- fouling and pre-treatment, and
- backflushing and cleaning.

Staging is of critical importance in large-scale dense membrane processes, i.e. reverse osmosis, nanofiltration and electrodialysis. Fouling is ubiquitous throughout the entire gamut of membrane technologies, but pretreatment to suppress or ameliorate fouling is only routinely practised in dense membrane processes. Backwashing is always carried out when the module design permits this, and cleaning is an essential part of membrane plant operation.

2.4.1 Staging

For most dense membrane processes the conversion of feed into product is limited either by the membrane area or the rate of extraction attainable by passage through a single module. It is for this reason that most reverse osmosis and electrodialysis technologies employ staging, the use of sequential stages to produce more product than that attainable by a single passage.
In reverse osmosis and nanofiltration even the largest membrane elements cannot normally achieve a recovery of much more than 20%, and the onset of concentration polarisation and the scaling this produces normally limits the conversion to well below this figure. It is therefore normal for them to be placed in series, with the retentate stream from one element being passed on to the feed stream of the next (Fig. 2.19). As many as eight or nine elements may be placed in a single module, and the resultant retentate flow exiting the module is then given by (from Equation (2.3)):

\[ Q_R = Q(1 - \Theta)^n \]  

(2.22)

where \( Q \) is the feed flow rate, \( \Theta \) the conversion per element and \( n \) the number of elements per module.

As the water flows along the length of a module the overall conversion is increased until there may come a point at which the element is running well below capacity. For example if, in Fig. 2.19, the conversion is 16% per element then the flow in stream 8 will be \((1 - 0.16)^4 \sim 0.5\) that of the feed. Under such
circumstances, it is advantageous to combine streams from modules operating in parallel in order to maximise utilisation of the membrane elements, such that an array of modules is produced (Fig. 2.20). This is known as staging, and is very common in the RO treatment of thin to light brackish waters where the retentate solution osmotic pressure does not become excessive through its concentration along the length of the module.

It is not necessarily feasible or desirable for the retentate to be staged in order to maximise the overall recovery. For high-salinity feedwaters, typically seawater where osmotic pressures in the region of tens of bar prevail, further concentration of these waters would demand uneconomically large operating pressures. For these and other waters permeate staging is employed, and the retentate from the second stage is returned to the feed of the first stage to produce a so-called twin pass system (Fig. 2.21). Twin pass systems also find use in high-purity water production.

Staging in electrodialysis differs from reverse osmosis in that further desalting of a solution that has already been desalinated by a maximum of 50% for a single passage through the stack, as constrained by the limiting condition imposed by depletion polarisation (Section 2.3.2), demands either increasing the volumetric flow, and/or reducing the current passed. Since the current passed is directly proportional to the total equivalent amount of ions transferred (Section 2.4.2), the simplest way of achieving additional desalination is by simply directing the product from the first passage through half the number of cell pairs in the second passage, thereby doubling the flow rate whilst leaving the current constant. This is known as hydraulic staging (Fig. 2.22). If further desalination is required, it may become expedient to reduce the current across the stack. This is referred to as electrical staging. It is normal to combine hydraulic with electrical staging to achieve reasonable desalination levels.

*Figure 2.21  Twin-pass RO system (permeate staged)*
2.4.2 Specific energy demand

Pressure-driven processes

The energy consumption of any pressure-driven process operating continuously is given by product of the pressure and flow, thus the total energy consumption per unit mass of permeate product, ignoring pumping efficiency, is given by:

\[ E = \frac{1}{\rho} \sum \Delta P + \frac{W_{el}}{M} \]  \hspace{1cm} (2.23)

where \( \rho \) is the permeate density, \( W_{el} \) the supplementary plant electrical energy consumption (e.g. for instrumentation and aeration or other turbulence promotion methods) and \( M \) the mass flow of permeate. \( \Sigma \Delta P \) represents the sum of the individual pressure changes, which can include the pressure drop across the membrane \( \Delta P_m \), as given by Equation (2.5), the concentrate–permeate osmotic pressure difference \( \Delta \Pi^* \), from Equation (2.9), the hydraulic losses \( \Delta P_{\text{losses}} \) associated with forcing the retentate through the membrane channels and the pressure derived from the various contributions to membrane fouling. In the case of reverse osmosis, the total pressure for an individual membrane module is given by:
\[ \Delta P_{\text{total}} = \mu J R \theta + Q_F (1 - \theta) \frac{dP}{dQ} + \frac{yRT \sum C_i}{1 - \theta} \]  

(2.24)

where \( Q_F \) is the feed flow rate to the module and fouling is ignored. In Equation (2.24), the pressure losses per unit flow through the membrane channels on the retentate side are represented by \( dP/dQ \). This factor can be estimated from the Hagen-Poiseulle equation, but is normally obtainable from the membrane supplier. An allowance can be made for fouling through an empirical correction factor. The rate of fouling in most reverse osmosis applications is normally low, since fouling is routinely suppressed by appropriate pretreatment based on chemical addition (Section 2.4.3).

In the case of filtration, the osmotic pressure term (the last term in Equation (2.24)) does not apply but the effects of cake formation and fouling on operating pressure cannot be ignored. Large-scale dead-end filtration plant operates with a backflush cycle actuated either at fixed intervals or when the membrane permeability has decreased to some pre-identified level. Since the backflush invariably fails to entirely recover the original membrane permeability, a more rigorous cleaning cycle, usually involving aggressive chemicals, is initiated again either at fixed intervals or when once the permeability of the backflushed membrane has declined to some other level. The calculation of energy consumption per unit volume must therefore incorporate the effects of downtime, specifically the energy consumption of the backflush and duration of the backflush and cleaning cycles (Section 4.3.5). Since backflushing is normally at 3–4 times the forward-flow flux, the energy expenditure per unit time is commensurately higher.

Dead-end filtration plant may be operated at either constant pressure or at constant flux (Fig. 2.23). In both cases trends in permeability decay over the backflush cycle, as reflected by flux decline at constant pressure operation (Fig. 2.23a) or pressure increase at constant flux operation (Fig. 2.23b), is exponential or pseudo-exponential, although the exact trend is dependent on the relative contributions of cake filtration, pore blocking and adsorption (Table 2.8). For filtration of solids forming an incompressible cake, on the other

\[ \begin{align*}
0 & \quad 5 & \quad 10 & \quad 15 & \quad 20 \\
0 & \quad 50 & \quad 100 & \quad 150 & \quad 200 & \quad 250 & \quad 300 \\
\text{Flux (L/MH)}
\end{align*} \]

\[ \begin{align*}
0 & \quad 2 & \quad 4 & \quad 6 & \quad 8 & \quad 10 & \quad 12 & \quad 14 \\
0 & \quad 200 & \quad 400 & \quad 600 & \quad 800 & \quad 1000 & \quad 1200 \\
\text{TMP (mbar)}
\end{align*} \]

Figure 2.23 Dead-end filtration: (a) constant pressure (declining flux over backflush cycle) and (b) constant flux (increasing pressure over backflush cycle)
hand, the trend is linear for constant flux operation. The trend in the residual membrane permeability, that is the permeability of the "permanently fouled" membrane as reflected in the trend in the minima for the backflush cycles in Fig. 2.22, also tends to be exponential or pseudo-exponential. Whether constant flux or constant pressure operation, the mean energy consumption relates to the mean membrane permeability over a cleaning cycle.

**Electrodialysis**

In electrodialysis the total power consumption is the product of the voltage and the current. Since the voltage relates directly to the current by the resistance according to Ohm's law, the specific energy demand is critically dependent upon the current and electrical resistance. The applied current \( i \) is directly proportional to the total equivalent quantity of counter-ions or co-ions extracted:

\[
i = \frac{QF\Delta C}{N\xi}
\]  

(2.25)

where \( Q \) is the flow through the stack, \( F \) is the Faraday constant, \( \Delta C \) is the change in the diluate concentration in eq m\(^{-3}\) from inlet to outlet. \( N \) is the number of cell pairs (the number of desalinating or concentrating cells in the stack) and \( \xi \) is the current efficiency (normally 85–95%). The energy demand is then given by:

\[
E = \frac{i^2R}{\Theta Q}
\]  

(2.26)

where \( R \) is the overall electrical resistance in ohms, and relates to the cell pair resistance \( R_{CP} \):

\[
R = R_{CP}N
\]  

(2.27)

If scaling can be overcome and the path length can be extended by staging the principal constraint placed on the recovery \( \Theta \) in Equation (2.26) is from electrodialysis reversal (see below). Because there is no osmotic pressure limitation in electrodialysis, and because the concentrate and diluate streams can be completely segregated, further efficiencies are obtained by recycling of the concentrate stream, which then decreases both the concentrate waste volume and the electrical resistance across the stack. For large-scale systems, recoveries in excess of 80% are normal.

The cell pair electrical resistance is given by:

\[
R_{CP} = R_{conc} + R_{dil} + R_{CEM} + R_{AEM}
\]  

(2.28)

where the subscripts conc and dil refer to the concentrate and diluate compartments and CEM and AEM refer to the cation and anion exchanging membranes. The concentrate resistance is normally no more than 20% of the
diluate resistance and the resistance of the ion exchange membranes, although determined to some extent by the salt concentration, is normally low – between 2 and 10 Ω cm² for most commercial ion exchange membranes in 0.5 eq m⁻³ NaCl (Strathmann, 1984). It therefore follows that the main contribution to the electrical resistance is from the diluate cell.

The specific energy demand is increased, by 5–10%, by electrodialysis reversal (EDR) operation. In this operational mode, which is very common in ED applications, the current is periodically reversed such that the concentrated stream becomes the diluate stream and vice versa. This adds to process complexity and reduces recovery but also virtually eliminates problems with scaling, since reversing the polarity of the electrodes produces a concomitant pH shift that suppresses the build-up of scale during the high-pH cathode electrode compartment.

**Ancillary mass transfer promotion**

Mass transfer can be promoted simply through changing the spacer material. Membrane spacers for ED technologies are either sheet flow, which promote plug flow of the water throughout the width of the channel, or tortuous path, which provide an extended path length and thus a greater linear flow velocity for the same volumetric throughput (Fig. 2.24). Tortuous path spacers yield higher Reynolds numbers but at the expense of a higher pressure drop across the stack. Most commercial ED systems are now based on sheet flow spacers.

Certain technologies use additional devices to promote mass transfer. Examples include the mechanically enhanced, high-shear processes (Section 2.1.4) which employ motors to increase the shear at or near the membrane surface. The additional energy consumption of such devices relates directly the shear applied, and a key part of the design of such devices is optimisation so as to impart the maximum shear for the minimum energy expenditure. As already stated (Section 2.1.4), coarse bubble aeration may also be employed to enhance mass transfer and in immersed membrane processes, such as the submerged membrane bioreactor (Stephenson et al., 2000). In such cases it represents a significant, sometimes the most significant, contribution to the overall energy demand and is dependent on the air flow rate, the nozzle diameter and the submersion depth. Other turbulence promotion devices, such as pulsed and vortex flow (Section 2.1.4) have yet to be commercialised.

![Electrodialysis spacers: (a) sheet flow (Eurodia) and (b) tortuous path (Ionics)](image.png)
2.4.3 Fouling and pretreatment

Pretreatment is required to suppress fouling and/or clogging of membranes, or else to remove chemically aggressive constituents such as chlorine. Clogging of membrane channels by solid matter can, to some extent, be controlled hydrodynamically, for a cross-flow process, or by the appropriate backflush regime for a dead-end process. It is only for certain filtration processes and/or specific duties, such as filtration of municipal or laundry wastewater, that pretreatment to remove gross solids may be critical. In these examples it is the filamentous matter that causes a problem, as it can form large aggregates which can become tangled with hollow fibres or otherwise clog the membrane channels and, for an immersed system, the aerators. Submerged membrane bioreactors for sewage treatment thus routinely employ fine screens and/or microstrainers to remove these materials.

Pretreatment of membrane filtration feedwaters can be analogous to that employed for depth filtration. An example is the use of coagulants to assist in the removal of natural organic matter (NOM) from upland surface waters. Precoagulation increases overall NOM removal, thereby reducing the propensity to form trihalomethanes, as well as producing a more permeable and less adherent filter cake (Judd and Hillis, 2001).

The requirement for pretreatment is determined by the feedwater concentration of:

- suspended solids, and colloidal matter in particular,
- scalants (sparingly soluble dissolved salts), and
- microorganisms and nutrients.

The physical manifestation of these are briefly described in Table 2.13. Appropriate pretreatment technology can normally only be identified through pilot plant trials or through reference to appropriate case studies. However, in

<table>
<thead>
<tr>
<th>Foulant</th>
<th>Symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silt/carbon fines</td>
<td>Element may be stained with brown or black material at the inlet and exhibit low permeate flow. Higher flow and very poor rejection may occur in later stages due to irreparable damage to the membrane by abrasive particles.</td>
</tr>
<tr>
<td>Carbonate scale, suspended silica</td>
<td>Usually on tap water or brackish water elements only. The element may be noticeably heavier than normal, and will exhibit low permeate flow and poor salt rejection.</td>
</tr>
<tr>
<td>Iron fouling</td>
<td>Rust colouring, possibly originating from iron pipework, at inlet of element. Element will exhibit low permeate flow and poor salt rejection.</td>
</tr>
<tr>
<td>Biological growth</td>
<td>Element may have strong odour, possible mould growth and will exhibit low permeate flow whilst maintaining a high salt rejection, which can increase with decreasing flux.</td>
</tr>
</tbody>
</table>
the case of pressure-driven dense membrane processes (reverse osmosis and nanofiltration), pretreatment is both critical and to some extent predictable from the feedwater quality – specifically from derived indices pertaining to colloidal particles and calcium carbonate scalant. Proscriptive methods for measuring fouling propensity are available (ASTM D4189, D3739, D4582, D4692, D4993), and in the following sections pretreatment requirements for reverse osmosis are discussed. A discussion of cleaning methods for fouled membranes follows in Section 2.4.4.

**Suspended solids**

Suspended solids in the feed can accumulate at the membrane surface to an extent dependent upon the degree of turbulence provided by the cross-flow. Gross suspended solids are readily removed by pre-filtration using cartridge filters (Section 2.1.4) which may or may not be preceded by depth (e.g. sand) filtration. Colloidal matter, derived from aluminium silicate (clays), iron colloids, organic materials, etc., is less readily removed. Colloids introduce problems in membrane processes only when coagulation of the suspended colloidal particles takes place in the membrane. Pretreatment is thus designed either to remove the colloidal particles or else stabilise them to prevent their coagulation during permeation.

The fouling nature of a water is best determined by the Fouling Index or Silt Density Index (SDI), which measure the rate at which a membrane’s pores plug. Standard test kits are available for determining SDI values. The test, a standard empirical test described in the reference literature (ASTM D4189), involves passing water at constant pressure (typically 2 bar) through a standard 0.45 μm-rated filter. The time $t_i$ taken to collect a given volume of water, typically 100 ml, from the clean filter is measured. Filtration is continued for a pre-set time ($t_t$), usually 15 minutes, and the time ($t_f$) taken to collect a second sample of the same volume is measured. The SDI is then given by:

$$SDI = \frac{100(1 - t_i/t_f)}{t_i}$$  \hspace{1cm} (2.29)

Suppliers of spiral wound RO and NF modules will normally specify an SDI value below 5. Waters of higher SDI values must be pretreated either chemically to stabilise the colloid, normally by addition of chemicals although occasionally softening can be used to stabilise the colloid by removing colloid-destabilising divalent species, or by some solid–liquid separation process to remove suspended material. Softening is unlikely to be economically viable on this basis alone, but in reducing the divalent ion level the scaling propensity of the water is also reduced which then allows operation at a higher conversion. Physical separation may demand pre-coagulation to increase the particle size and ensure retention of the suspended solids by the filter. For some applications where the feedwater contains natural organic matter the RO process may be preceded by fine microfiltration or even ultrafiltration.
Membrane technology  61

Scalants and scaling indices

Scalants are low-solubility salts whose precipitation onto the membrane is promoted by the conversion of water into permeate and further encouraged in pressure-driven dense membrane processes both by concentration polarisation and the pH shift produced by carbon dioxide permeation. The scale formed can reduce the membrane permeability and permselectivity. As with colloidal and particulate fouling, scaling is also a problem in membrane filtration processes. Any water containing calcium carbonate close to or beyond its thermodynamic saturation limit, as is the case for many dairy and pharmaceutical effluents, can produce calcite (the most common crystal form of calcium carbonate) at the membrane surface.

Scale formation propensity is usually apparent from chemical thermodynamics, and specifically the solubility product $K_s$ (Table 2.14), although it can never be unequivocally predicted. The solubility product represents the maximum value of the product of the molar concentrations of the two component ions of the salt. If the solubility is exceeded then the salt will precipitate. The general rule of thumb to avoid precipitation is that the ionic product should not exceed 80% of the solubility product. The appropriate constants for thermodynamic equilibria appropriate to some of the more common scalants, such as salts of the divalent alkaline earth elements of magnesium, calcium and barium, are normally included in CAD packages for designing RO arrays (Section 4.1). The thermodynamic relationships include, in the case of calcium carbonate formation, data pertaining to hydrolysis. The significance of this is outlined below.

Calcium carbonate is very insoluble in water and readily precipitates to form a scale on pipework, heat transfer surfaces and membranes. The equilibrium constant for the dissolution reaction is represented by:

$$K_s = [Ca^{2+}][CO_3^{2-}]$$

and so:

$$[CO_3^{2-}] = K_s/[Ca^{2+}]$$

When carbon dioxide dissolves in water it forms carbonic acid, which dissociates producing acid and bicarbonate ions thus:

$$CO_2(\text{dissolved}) + H_2O\rightarrow H^+ + HCO_3^-$$

This is the origin of the pH shift in reverse osmosis. Because the membrane allows free passage of carbon dioxide, the CO$_2$/HCO$_3^-$ ratio in the permeate is high and that of the retentate low. Since, according to Equation (2.32), the acid (i.e. H$^+$) concentration relates directly to the CO$_2$/HCO$_3^-$ ratio, the retentate pH is correspondingly high whilst that of the permeate is low.

Bicarbonate ions further dissociate to carbonate:

$$HCO_3^- + H^+ \rightarrow CO_3^{2-}$$
### Table 2.14 Solubility products for some common scalants at 20°C

<table>
<thead>
<tr>
<th>Salt</th>
<th>Formula</th>
<th>Solubility product, $K_s$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminium hydroxide</td>
<td>Al(OH)$_3$</td>
<td>$8.5 \times 10^{-21}$</td>
</tr>
<tr>
<td>Barium sulphate</td>
<td>BaSO$_4$</td>
<td>$9.2 \times 10^{-1}$</td>
</tr>
<tr>
<td>Calcium carbonate</td>
<td>CaCO$_3$</td>
<td>$4.8 \times 10^{-9}$</td>
</tr>
<tr>
<td>Calcium fluoride</td>
<td>CaF$_2$</td>
<td>$3.2 \times 10^{-11}$</td>
</tr>
<tr>
<td>Calcium hydrogen phosphate</td>
<td>CaHPO$_4$</td>
<td>$2.0 \times 10^{-7}$</td>
</tr>
<tr>
<td>Calcium hydroxide</td>
<td>Ca(OH)$_2$</td>
<td>$8 \times 10^{-6}$</td>
</tr>
<tr>
<td>Calcium sulphate</td>
<td>CaSO$_4$</td>
<td>$2.3 \times 10^{-4}$</td>
</tr>
<tr>
<td>Cadmium sulphide</td>
<td>CdS</td>
<td>$1.4 \times 10^{-28}$</td>
</tr>
<tr>
<td>Cobalt sulphide</td>
<td>CoS</td>
<td>$3.0 \times 10^{-26}$</td>
</tr>
<tr>
<td>Chromium hydroxide</td>
<td>Cr(OH)$_3$</td>
<td>$2.9 \times 10^{-29}$</td>
</tr>
<tr>
<td>Copper sulphide</td>
<td>CuS</td>
<td>$1.0 \times 10^{-44}$</td>
</tr>
<tr>
<td>Dolomite</td>
<td>CaMg(CO$_3$)$_2$</td>
<td>$6.8 \times 10^{-18}$</td>
</tr>
<tr>
<td>Ferrous hydroxide</td>
<td>Fe(OH)$_2$</td>
<td>$4.8 \times 10^{-16}$</td>
</tr>
<tr>
<td>Ferric hydroxide</td>
<td>Fe(OH)$_3$</td>
<td>$3.8 \times 10^{-18}$</td>
</tr>
<tr>
<td>Ferrous sulphide</td>
<td>FeS</td>
<td>$4.0 \times 10^{-19}$</td>
</tr>
<tr>
<td>Lead sulphide</td>
<td>PbS</td>
<td>$5.0 \times 10^{-29}$</td>
</tr>
<tr>
<td>Manganese hydroxide</td>
<td>Mn(OH)$_2$</td>
<td>$4.0 \times 10^{-14}$</td>
</tr>
<tr>
<td>Manganese sulphide</td>
<td>MnS</td>
<td>$1.4 \times 10^{-15}$</td>
</tr>
<tr>
<td>Magnesium ammonia phosphate</td>
<td>MgNH$_4$PO$_4$</td>
<td>$2.2 \times 10^{-11}$</td>
</tr>
<tr>
<td>Magnesium carbonate</td>
<td>MgCO$_3$</td>
<td>$1.0 \times 10^{-5}$</td>
</tr>
<tr>
<td>Magnesium hydroxide</td>
<td>Mg(OH)$_2$</td>
<td>$3.4 \times 10^{-11}$</td>
</tr>
<tr>
<td>Mercuric bromide</td>
<td>Hg$_2$Br$_2$</td>
<td>$5.2 \times 10^{-21}$</td>
</tr>
<tr>
<td>Mercuric chloride</td>
<td>Hg$_2$Cl$_2$</td>
<td>$3.5 \times 10^{-18}$</td>
</tr>
<tr>
<td>Mercuric sulphide</td>
<td>Hg$_2$S</td>
<td>$1.0 \times 10^{-45}$</td>
</tr>
<tr>
<td>Nickel hydroxide</td>
<td>Ni(OH)$_2$</td>
<td>$8.7 \times 10^{-19}$</td>
</tr>
<tr>
<td>Nickel sulphide</td>
<td>AgBr</td>
<td>$7.7 \times 10^{-13}$</td>
</tr>
<tr>
<td>Silver chloride</td>
<td>AgCl</td>
<td>$1.5 \times 10^{-10}$</td>
</tr>
<tr>
<td>Silver sulphide</td>
<td>Ag$_2$S</td>
<td>$1.6 \times 10^{-49}$</td>
</tr>
<tr>
<td>Zinc hydroxide</td>
<td>Zn(OH)$_2$</td>
<td>$1.0 \times 10^{-17}$</td>
</tr>
<tr>
<td>Zinc sulphide</td>
<td>ZnS</td>
<td>$1.0 \times 10^{-23}$</td>
</tr>
</tbody>
</table>

For which the equilibrium constant is:

$$K_2 = \frac{[H^+][CO_3^{2-}]}{[HCO_3^-]}$$

and so:

$$[CO_3^{2-}] = K_2[HCO_3^-]/[H^+]$$

Equations (2.31) and (2.35) can be combined to give:

$$[H^+] = K_2[Ca^{2-}][HCO_3^-]/K_s$$

Thus:

$$pH_s = (pK_2 - pK_s) + pCa + pAlk$$
where Alk represents the molar bicarbonate concentration or alkalinity. In this equation the term pH_s represents the pH at which the water is in equilibrium with calcium carbonate, and is thus sometimes referred to as the saturation pH. The value of (pK_2 - pK_s) is dependent on temperature and ionic strength.

The Langelier Saturation Index is defined as:

\[
\text{LSI} = \text{pH} - \text{pH}_s
\]  

where pH is the actual measured pH of the water and pH_s is the calculated value. The LSI effectively measures the degree to which the water is either supersaturated or undersaturated with calcium carbonate, and thus its propensity for forming scale (ASTM D3739). A negative LSI indicates a corrosive water that will dissolve calcium carbonate scale and a positive LSI indicates a scale-forming one. The LSI is widely used and there are several nomograms available for rapid calculation (Fig. 2.25).

Whilst the LSI is a good predictor of scaling it is a bit unreliable in the range -1 to +1 and gives no indication of the relative scaling potential. The Ryznar Stability Index gives more quantifiable results and is defined as follows:

![Langelier Scaling Index nomogram](Figure 2.25)
Membranes for Industrial Wastewater Recovery and Re-use

RSI = 2pHs − pH (2.39)

An RSI of 7 indicates a water more or less at equilibrium. As the value falls the water becomes more scaling and as it rises the water becomes more corrosive. The Langelier and Ryznar saturation indices can be used to predict scaling in most waters but they become unreliable when the total dissolved solids content approaches 5000 mg l⁻¹. Above this level the Stiff and Davis index (ASTM D4582) or a similar method must be adopted.

Not all scale-forming compounds have a pH-dependent deposition. Some scalants have simple chemistry, analogous to that given in Equation (2.30):

\[ M^{2+} + \gamma^{\text{MY}} \]  

or

\[ M^{2+} + Z^{\text{MY}^Z} \]  

Scalants having the chemistry represented by Equation (2.39) are mainly the sulphates of magnesium, barium and strontium (ASTM D4692).

The simplest means of preventing scale formation in RO systems is to operate at a conversion sufficiently low that the reject stream is not so concentrated that solubility problems are encountered. This, of course, has economic implications for the operation of the plant that may be unacceptable, and in such cases some form of scale prevention must be used. Where calcium carbonate or some other hydrolysable scalant is the main problem, i.e. those salts containing hydroxide (OH⁻) and carbonate (CO₃²⁻) which therefore have associated pH-dependent solubility due to hydrolysis reactions (i.e. reaction with acid, H⁺) of these anions, it is often possible to adjust the LSI by acid dosing. This converts bicarbonate into carbon dioxide which can be removed by degassing of either the feed or the permeate. Depending on whether hydrochloric or sulphuric acid is used there will be an increase in sulphate or chloride concentration. Increasing the sulphate may give rise to calcium sulphate precipitation.

Scale-inhibiting chemicals can be used to delay precipitation of some salts by interfering with the crystallization process forming microcrystals which do not cause fouling and which will not show significant agglomeration at least until the concentrate stream has left the RO unit. Most of the commercially available chemicals work well on calcium carbonate, provided that the LSI in the concentrate stream is less than +2, but less well on other salts. The most established scale inhibitors are termed “glassy polyphosphates”, typically sodium hexametaphosphate (Calgon). These work by absorbing into the nanoscopic protonuclei forming during the incipient stages of precipitation and destabilising the subsequent crystal nuclei. They are not the most effective chemicals for the application and revert quite quickly in solution to orthophosphate, producing calcium phosphate sludge which can cause blocking of separators and small bore pipes. Since the development of Calgon, more
effective threshold chemicals, including phosphonates, have become available. Polycarboxylic acids, such as polyacrylates and polymalonates, operate by blocking crystal growth sites preventing the growth of nuclei into crystals whilst chelating chemicals react with potentially insoluble cations like calcium to form a soluble complex. Dosage rates are typically of the order of 5–10 mg l\(^{-1}\) and, whilst some indication of the efficacy of these reagents may arise from the scale inhibition mechanism, it is generally the case that only pilot trials yield reliable information as to their suitability for a particular duty.

Where scale-inhibiting chemicals are unable to cope with the concentrations involved, and where chemical dosing is to be avoided for some reason, then pretreatment of the water is necessary to remove the scale-forming salts. This may be by sodium cycle ion exchange softening, ion exchange dealkalisation, lime or lime-soda softening or even a nanofiltration process if divalent ions are to be selectively removed. Some scalants are particularly recalcitrant. Dissolved (or "active") silica, evaluated using ASTM D4993, is not readily removed by pretreatment and there are proprietary reagents that have been developed specifically to inhibit its precipitation. However, modern RO membranes are reasonably tolerant, and raw environmental waters rarely require more than filtration and dosing as pretreatment for membrane permeation.

**Microorganisms and nutrients**

Bacteria are ubiquitous and thrive in the high surface area environment of a reverse osmosis membrane element, where they form biofilms. They are naturally transported towards the membrane surface under the force of the permeate flow, and are supplied with nutrients both from the feedwater and, under some circumstances, the pretreatment chemicals. Reports of the latter, as discussed by Flemming (1992), include flocculants (Graham *et al.*, 1989), phosphorus-containing scale inhibition chemicals (Ahmed and Alansari, 1989), sodium thiosulphate, as used for quenching chlorine (Winters and Isquith, 1979), and even chlorine itself, which was found to degrade organic matter sufficiently for it to become biologically assimilable (Applegate *et al.*, 1986). Although they can form substantial biofilm layers in potable and wastewater applications, bacteria require only an extremely low nutrient load to survive in biofilms and can exist even in ultrapure water systems, where their management becomes particularly vexing. Biofilm formation is thus unavoidable in most membrane operations, and it is likely that all membrane fouling is associated with biofilm formation, although the biofilm itself is not necessarily onerous.

Biofilm formation results from the rapid formation of an organic film, normally within the first few minutes of operation, followed by microbial adhesion and further entrapment of dissolved and suspended solid matter which co-deposits with the microorganisms. The conditioning film can be existing organic matter from the feedwater, such as NOM, or microbial products, namely extracellular polymeric substances (EPS). The rate of microbial deposition and the overall biofilm thickness depend on hydrodynamics, with biofilm thickness apparently decreasing with increasing turbulence (Ridgeway, 1988). Other key factors
include the membrane material and feedwater bacterial and nutrient concentration (Characklis and Marshall. 1990).

The principle effect of biofilm formation on membrane process performance is to reduce membrane permeability. In the case of cellulosic membranes the bacteria may ingest the membrane itself, causing irreversible degradation and significant loss of salt rejection. Since the elimination of biofouling is only possible through complete removal of microorganisms, the only convenient effective pretreatment is dosing with a liquid biocide. However, the most effective of these (chlorine and its compounds) are oxidative and will degrade most reverse osmosis membranes, the principal exception to this being cellulosic materials. UV irradiation, and even pre-sterilisation, are of limited efficacy since they generally are unable to completely prevent biofilm formation and cannot act directly upon the biofilm once it has formed. The main emphasis, therefore, is on control of the biofilm (where possible), and on its periodic removal through an appropriate cleaning protocol.

**Organic matter**
There are essentially three key categories of organic foulants:

- proteins,
- carbohydrates, and
- fats, oils and grease (FOG).

As with biological matter, pretreatment to remove organic matter is rarely feasible, and it is more usual to adopt an appropriate backflush and, in particular, cleaning strategy so as to ameliorate the worst effects of organic fouling. Proteins and carbohydrates form a part of the extracellular polymeric substances (EPS), as well as natural organic matter (NOM). Proteins may be colloidal or dissolved, and are least soluble at their isoelectric point, which tends to arise at pH values of 4–5. Their removal is therefore most effective at extremes of pH. Carbohydrates include starches, polysaccharides, and fibrous and pectin materials. Fouling by these materials is very sensitive to flux, and their removal sensitive to the precise cleaning protocol adopted. FOG forms part of the suspended matter, and is particularly problematic since these substances generally have a high affinity for the more hydrophobic membranes, and polysulphone in particular, and are not readily removed by normal backflushing and cleaning methods. Their effective removal is normally only achievable at high temperatures and/or by the use of organic solvents. Organic solvents are only an option for ceramic or highly inert polymeric membranes (Section 2.1.3).

A summary of foulant impacts and pretreatment options for reverse osmosis are listed in Table 2.15.

### 2.4.4 Backflushing and cleaning

An essential distinction must be made between the intermittent backflush cycle, in which the fouled membrane is physically cleaned by flowing the product
Table 2.15  Common foulants and their pretreatment, reverse osmosis

<table>
<thead>
<tr>
<th>Fouulant</th>
<th>Effect on performance</th>
<th>Pretreatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suspended solids</td>
<td>Decrease in permeability, normally in first stage</td>
<td>&lt; 2 mg l(^{-1}); 5 μm cartridge filter;&lt;br&gt; &gt; 2 mg l(^{-1}); multimedia deep-bed filter&lt;br&gt; Cross-flow microfiltration</td>
</tr>
<tr>
<td>Organic and inorganic colloidal matter, SDI &gt; 5</td>
<td>As above</td>
<td>Ultrafiltration; Precoat filtration; Multi media deep bed filtration; In-line coagulation and filtration; Softening</td>
</tr>
<tr>
<td>Alkaline scales: CaCO₃</td>
<td>Decrease in salt rejection and slight decrease in permeability, normally in final stage</td>
<td>LSI &lt; 2; threshold inhibitor dosing; LSI &gt; 2; acid addition; LSI &gt; 2; ion exchange or lime-soda softening</td>
</tr>
<tr>
<td>Active silica</td>
<td>As above</td>
<td>Lime-soda softening with MgO; Alkali dosing; Heat</td>
</tr>
<tr>
<td>Permanent hardness</td>
<td>As above</td>
<td>Scale inhibitor dosing, ion exchange softening, lime-soda softening; Threshold inhibitors</td>
</tr>
<tr>
<td>Sulphate scales, e.g. CaSO₄</td>
<td>As above</td>
<td>Aeration and filtration, dissolved air flotation (DAF), catalytic oxidation (greensand filtration), chemical oxidation and filtration</td>
</tr>
<tr>
<td>Metal oxide (iron and manganese)</td>
<td>As above</td>
<td>Activated carbon, chemical reducing agent</td>
</tr>
<tr>
<td>Free chlorine</td>
<td>Permanent loss of rejection due to degradation of membrane*</td>
<td>Adsorption (GAC), flotation</td>
</tr>
<tr>
<td>Dissolved organics, e.g., NOM, EPS, anionic detergents</td>
<td>Rapid decrease in permeability</td>
<td>Chlorination/dechlorination, UV (+ microfiltration)</td>
</tr>
<tr>
<td>Biological matter</td>
<td>Decrease in permeability</td>
<td></td>
</tr>
</tbody>
</table>

*Non-chlorine-resistant materials.

Permeate water in the reverse direction through the membrane, and the cleaning cycle in which the membrane is invariably chemically treated for a more extended period. Whilst both backflushing and cleaning arc routinely and advantageously employed to recover some of the flux, nonetheless incur a cost penalty through:

- the energy consumption,
- the loss of product water,
- the downtime, and
- the use of hazardous chemicals.

The energy consumption relates to the hydraulic resistance, as discussed in Section 2.4.2. The loss of product water arises because of its use for both backflushing and cleaning, and can be significant. For example, a 20 s backflush
every 12 minutes, as might be appropriate for a dead-end filtration of a municipal wastewater, incurs a loss of 8% of the product water if the backflush flow is three times that of the forward flow. The cleaning cycle employs less product water and generally a shorter percentage downtime but demands the use of acidic, caustic and/or oxidative chemicals and so produces extra transport and handling problems and generates a waste stream which may not be so readily disposed of as the backwash water. Cleaning, of course, also incurs the additional cost of both the cleaning reagent(s) and system.

Cleaning chemicals generally fall into one of seven categories, with some overlap between them:

- Acids
- Alkalis
- Chelants
- Enzymes
- Surfactants
- Biocides
- Oxidants

A wide variety of proprietary formulated membrane cleaning products have been developed and marketed mainly by the membrane suppliers but also by other companies, with the number of products proliferating over the last decade or so. Notwithstanding these commercial developments, many membrane plant operators still tend to employ conventional generic cleaning chemicals and formulations (Table 2.16). It is also the case that cleaning solutions tend to be employed on a reactive rather than routine preventative basis. It is generally the case that employing cleaning once fouling of the membrane has become severe is less effective than routine maintenance cleaning which removes incipient fouling. Also, in many cases no single cleaning reagent will recover the flux entirely, and it is often advantageous to use a combination of cleaning reagents in sequence.

Table 2.16 Common foulants and appropriate cleaning solutions, reverse osmosis

<table>
<thead>
<tr>
<th>Foulant</th>
<th>Cleaning solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium carbonate</td>
<td>Hydrochloric acid pH 4</td>
</tr>
<tr>
<td>Calcium, barium</td>
<td>2% citric acid ammoniated to pH 4</td>
</tr>
<tr>
<td>sulphate</td>
<td>2% citric acid ammoniated to pH 8</td>
</tr>
<tr>
<td>Metal oxides</td>
<td>EDTA sodium salt, pH 7–8</td>
</tr>
<tr>
<td>Silica</td>
<td>2% citric acid ammoniated to pH 4</td>
</tr>
<tr>
<td>Organics</td>
<td>1% sodium hyposulphite</td>
</tr>
<tr>
<td>Biological fouling</td>
<td>Caustic soda pH 11</td>
</tr>
<tr>
<td></td>
<td>Caustic soda pH 11</td>
</tr>
<tr>
<td></td>
<td>Alkaline detergent</td>
</tr>
<tr>
<td></td>
<td>1% formaldehyde</td>
</tr>
<tr>
<td></td>
<td>Hypochlorite</td>
</tr>
</tbody>
</table>
Acid cleaners are generally used for inorganic scale removal, principally calcium carbonate and ferric hydroxide. Reagents employed include conventional low-cost inorganic acids (sulphuric, hydrochloric, nitric and, less commonly, phosphoric) and organic acids (citric, oxalic, acetic and sulphamic). The stronger, in terms of acid strength per equivalent concentration, inorganic acids are correspondingly more effective that the weaker organic salts but are also more corrosive and potentially more damaging to the membrane element, module or plant pipework. Careful pH control is therefore necessary, with pH values above 2 being maintained for most applications. Citric acid solutions, which have historically been employed for scale removal, must be amminated and adjusted to a pH of 4 to avoid complexation with the ferrous ion, since this can precipitate and blind the membrane. Oxalic acid is particularly effective against iron, but forms a calcium salt of limited solubility. Some organic acids act through chelation, i.e. through the formation of stable complexes with multivalent metal cations. The most commonly used chelants are ethylene diamine tetracetic acid (EDTA), phosphonocarboxylic acid, gluconic acid, citric acid and various polymers that have a chelating ability. Of these EDTA, as a hexadentate ligand, is significantly the stronger but also has adverse health effects.

Alkaline cleaners, primarily caustic soda, are employed for the removal of organic acid foulants, including proteins, and silica. The solution may need buffering with bicarbonate or phosphate to maintain the pH, however, and the addition of surfactants can improve the detergency action of the alkaline solution. It is common to combine caustic with acid cleaning and other cleaning chemicals, applied sequentially, to maximise the flux recovery. Surfactants themselves can be non-ionic (e.g. ethylene oxide), anionic (soaps and alkylsulphonates) or cationic (quaternary ammonium compounds or QACs). QACs, which are also bacteriostatic (i.e. prevent the growth of bacteria), cannot be used with membranes with a high surface concentration of carboxylate ions, such as polyamide thin-film composites, due to irreversible complexation. Non-ionic surfactants may also exhibit this behaviour, though to a lesser extent.

Enzymic cleaners are effective against many biodegradable organic foulants, such as proteins, and are chemically non-aggressive. They are, however, also slow acting and expensive, and so are generally only employed when other cleaning chemicals have failed. Since they are also proteins in themselves, an additional alkaline chemical clean is required to remove any residue.

The most commonly used biocide is chlorine, although aromatic polyamide membranes are degraded by chlorine and other oxidants. Generally chlorine levels of around 10 ppm are sufficient for routine sanitisation following a chemical clean, but concentrations of 100 ppm may be used for specific bactericidal duties such as pyrogen removal. Chlorine is generally employed as a sodium hypochlorite solution, adjusted to a pH of 10–11 to reduce its chemical aggressiveness, though this also reduces its bactericidal potency. A 2–3% formaldehyde solution can also be used for sanitisation, and this reagent is also used at lower concentrations (0.5–1%) for storing membranes as an alternative to the more potent hypochlorite. However, it must be removed by thorough
rinsing prior to membrane operation due to its propensity for complexation with proteins. Other chemical bactericidal reagents that have been employed, and may be included in proprietary formulations, include QACs, acidified metabisulphite, hydrogen peroxide and peroxyacetic acid, the latter being generated from hydrogen peroxide and peracetic acid. Some membranes, notably fluorinated polymers and ceramic membranes, are steam sterilisable.

Most membrane plant is designed to allow cleaning in place (CIP). Fouled cross-flow membranes systems are cleaned by recirculating a cleaning solution around the concentrate side of the module by means of a small tank and circulating pump. Each membrane module, or bank of modules if an entire bank is to be cleaned at once, is normally provided with cleaning connections to allow this. The connections are usually three-way L-port ball valves so that connecting the cleaning solution circulation system automatically isolates the module(s) from the high-pressure feed system. In dead-end systems CIP entails pumping the cleaning solution in the reverse direction, i.e. from the permeate to the retentate side. Submerged systems can also be cleaned ex situ by placing the entire module in a tank containing the cleaning fluid.

The cleaning tank for an RO plant normally has a working volume equal to the volume of the (empty) pressure tube(s) to be cleaned plus the volume of the associated pipework. Tanks are typically of GRP or polypropylene, depending on the cleaning chemical being used, and fitted with an electrically powered stirrer. The circulation pump is normally rated to deliver a flow velocity of 0.06–0.08 m s$^{-1}$ at a pressure 1.2–4 bar, the exact pressure being dependent upon the number of elements in the module. Precise recommended cleaning protocols vary, with some suppliers recommending a 15–30 minute soak. It is normally desirable for this stream to be fitted with a 5 µm filter upstream of the pressure vessels. Disinfection can be carried out in the same way but with a higher pressure (up to about 10 bar) to ensure that the disinfecting solution penetrates to the permeate side of the membrane.

References


ASTM D4189, Standard test method for Silt Density Index (SDI) of water.

ASTM D4582. Standard practice for calculation and adjustment of the Stiff and Davis stability index for reverse osmosis.
ASTM D4692, Standard practice for calculation and adjustment of sulfate scaling salts (CaSO₄, SrSO₄ and BaSO₄) for reverse osmosis.

ASTM D4993, Standard practice for calculation and adjustment of silica scaling for reverse osmosis.


Chapter 3

Industrial waters

3.1 The power industry
Audrey Levine
University of South Florida

3.2 The pulp and paper industry
Jutta Nuortila-Jokinen, Mika Mänttäri and Marianne Nyström
Lappeenranta University of Technology

3.3 The textile industry
Simon Judd
School of Water Sciences, Cranfield University

3.4 The beverage industry
Mike Carney
Vale Water Services Ltd

3.5 Pure waters in the pharmaceutical industry
John Hutcheson
Christ Kennicott Water Technology Ltd
3.1 The power industry: water requirements for power generation and cooling water

3.1.1 Water demand by the power industry

In industrialized nations significant quantities of water are needed for power generation and cooling applications, greatly exceeding the quantities of water used for all other purposes (Fig. 3.1). For example, in the USA thermoelectric power generation accounts for about 47% of the $5.5 \times 10^{11}$ m$^3$ of water that is consumed annually (Solley et al., 1998). A major water use is for condenser and reactor cooling in steam–electric power plants. For power plants equipped with wet towers for steam condensation, the evaporation of water in the cooling tower can account for up to two-thirds of the total plant water use. Other applications for cooling water include air conditioning systems, food and beverage manufacture, meat and dairy industries, chemical manufacturers, and petroleum refineries (Asano and Levine, 1998).

On a global basis, it has been estimated that about 10 000 GW of new electrical generating capacity will be required by 2050 (Powicki, 2002). Fresh water is an essential resource for the production of electricity, as it is needed for high-purity steam, condensate cooling, and miscellaneous operations including dust suppression. While the net water requirements depend on the distribution of energy sources (nuclear, fossil fuel, waste-to-energy, hydropower, solar, wind, etc.), it is obvious that energy production is dependent on water availability. In addition to projected increases in energy and associated water requirements, liquid discharges from industrial water users are subjected to increasingly stringent requirements in many locations. In the future, it is likely that many thermoelectric power generation facilities will be required to comply with so-called zero liquid discharge (ZLD) requirements to meet long-term watershed

![Figure 3.1 Water use in the USA (data from Solley et al., 1998)](image)
protection goals, particularly in environmentally sensitive locations (Crook et al., 1994; Puckorius, 1997). It is thus important to develop more efficient approaches to management of water in the thermoelectric power industry.

Over the past decade, recurring water shortages coupled with more rigorous water quality requirements have prompted a re-examination of current cooling water practices. Water availability poses challenges to the siting of new power production facilities. In addition, a growing water resource management trend worldwide is to prioritise the use of water based on water quality and availability. Where appropriate, the emphasis is on preserving the highest quality water sources for potable water applications by using secondary sources for applications that have less stringent water quality requirements. As is evident from Fig. 3.1, efforts to reduce the quantity of water used for thermoelectric power generation can have a direct impact on balancing the distribution of water resources. This section aims to discuss the potential for using reclaimed water as a water source for cooling operations. A description of cooling water applications is given, water quality requirements are discussed and treatment alternatives presented. Some existing examples of cooling operations using reclaimed water are also given.

3.7.2 Overview of cooling water systems

Thermoelectric power plants typically derive energy from heat-generating fuels, such as nuclear fuel, coal, oil, natural gas, municipal solid waste, or refuse-derived fuel. The heat is used to produce steam in a boiler. The steam passes through a turbine and is then condensed back to water and pumped back to the boiler to repeat the cycle. The turbine drives a generator that produces electricity. Steam turbines extract power from steam as it passes from high-pressure and high-temperature conditions at the inlet to low-pressure and low-temperature conditions at the outlet. The outlet pressure is a function of the temperature of the cooling water used to absorb heat and reject it from the condenser (Asano et al., 1988; Burger, 1979; Kemmer, 1988).

The water requirements for power plants are based on providing water for cooling and water as a source of steam in boilers. Typically, higher quality water is needed for boiler feed water as compared to cooling water. In general, cooling systems function to remove excess heat from heat sources. A cooling tower acts as a heat exchanger by driving ambient air through falling water, causing some of the warmed water to evaporate, thereby dissipating heat and providing cooling. The cooler water is then circulated back to the equipment that needs cooling. In thermoelectric power plants, cooling water is used to remove heat from the turbine exhaust and condense it back to liquid prior to its return to the steam generator.

Cooling tower operations consist of evaporative condensation and exchange of heat. Latent heat of vapourisation is released from the air and water mixture, resulting in evaporation of water at a rate dependent on the temperature and pressure. The amount of heat consumed is about 2.3 kJ g⁻¹ of water evaporated. For each 5.5°C of cooling, about 1.2% of the water is released from the cooling
tower to the atmosphere as hot moist vapour. Evaporative cooling systems provide air flow either through natural draft or mechanical draft systems. Typically, natural draft systems are hyperbolic towers (> 50 m high) with air flow rates of about 2000 m³ min⁻¹ (Kemmer, 1988; Puckorius and Hess, 1991; State of California, 1980). Density gradients between the less dense heated air and the more dense cool ambient air result in positive air flow.

Mechanical draft systems employ fans to move air through the towers either through forced draft (fans at the bottom of the tower) or induced draft (fans at the top of the tower to pull air up through the tower). Rotating fans are used to pull cool, dry air into the tower at a rate of about 600–750 m³ air per m³ recirculating water. Warmed heat exchanger water is pumped to the top of the tower where it descends through the rising air stream. Packing material inside the tower breaks up the water flow to improve the contact efficiency between the air and water. A fraction of the water evaporates and leaves the tower as vapour. Mechanical draft cooling towers (Fig. 3.2) tend to have vapour plumes that are about 30% shorter than those produced by natural draft towers, thus reducing the need for plume abatement.

A variety of types of cooling systems are used, depending on the application and water availability (Table 3.1). Cooling water can be used once and returned to its original source (once-through cooling) or reused in an open recirculating

Figure 3.2  Vapour plume from cooling system at McKay Bay waste-to-energy facility, Tampa, Florida
Table 3.1  Comparison of water use in cooling systems (Asano et al., 1988; Blackson and Moreland, 1988; Burger, 1979; Kemmer, 1988; State of California, 1980; Tay and Chui, 1991)

<table>
<thead>
<tr>
<th>Type</th>
<th>Application</th>
<th>Water source</th>
<th>Water use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Once-through</td>
<td>Large power facilities or refineries adjacent to a water source</td>
<td>Fresh, saline, or reclaimed water</td>
<td>75–225 m$^3$ MWh$^{-1}$.</td>
</tr>
<tr>
<td>non-contact cooling</td>
<td></td>
<td>Evaporative consumption: 0.5–2%</td>
<td>Depends on process</td>
</tr>
<tr>
<td>Direct contact cooling</td>
<td>Processing of inert material when chemical reaction quenching needed, such as in primary metals industry</td>
<td>Freshwater; degree of purification related to process conditions</td>
<td></td>
</tr>
<tr>
<td>Non-contact, open, recirculating cooling</td>
<td>Widely used where water resources are limited and alternative water sources such as reclaimed water are available (such as inland regions)</td>
<td>Fresh, saline, or reclaimed water</td>
<td>0.5–4 m$^3$ MWh$^{-1}$. Evaporative consumption: 80–95%</td>
</tr>
<tr>
<td>Closed recirculating system</td>
<td>Water circulated in closed loop with negligible evaporation or exposure to atmosphere; Used for critical cooling applications</td>
<td>Ultrapure water</td>
<td>Water velocities: 1–1.5 m s$^{-1}$ Evaporative consumption: &lt; 5%</td>
</tr>
</tbody>
</table>

cooling system where process heat from the warmed water is transferred to the air via evaporation so that the water can be reused. Process cooling is accomplished through indirect contact of the process streams and cooling water in heat exchangers. Closed recirculating systems consume minimal water, but require higher purity water to prevent operational problems. Therefore, due to the economics of providing supplemental water treatment, closed recirculating systems are only used in cases where cooling efficiency is critical, such as in steel production and microelectronics. Currently, on a global basis, the majority of thermoelectric power plants rely on either once-through cooling or open recirculating cooling systems. The degree of water treatment required for cooling water applications depends on the system. Typically, minimal treatment is required for once-through cooling, whereas scale and corrosion control are needed for open recirculating systems. Closed recirculating systems require control of salts and minerals, similar to requirements for boiler operation.

In once-through processes, heat is transferred to water which is then returned to its source. For once-through cooling systems, the cooling water inlet temperature is the temperature of the source water, whereas the outlet temperature is 5–25°C higher than the inlet. Thus discharge of this water can introduce temperature gradients within the receiving water if inadequate mixing is available. Recirculating systems transfer the heat from the warmed water to the air through evaporation; therefore, outlet temperature and inlet temperatures are similar. As shown in Table 3.1, recirculating systems require less than 10% of the quantity of water used in once-through cooling systems. The reduction in water use has an impact upon the volume of water discharged from the cooling system and results in a net decrease in the quantity of thermal discharges.
3.1.3 Overview of boiler feedwater systems

An integral component of thermoelectric power production is the use of boilers to vaporise water into high-purity steam that is used to drive turbines. The operating pressure of a boiler influences the heat transfer rates, with higher pressures yielding rates of over 21 kJ m⁻² h⁻¹. Materials used for boilers include mild steel, copper alloys (admiralty brass or copper-nickel), or stainless steel (Kemmer, 1988). Cooling water either flows through the tubes (tubeside cooling) or through the shell and around the tubes. The two major types of boiler designs that are in widespread use include firetube and watertube systems. The firetube boilers tend to be used in boilers that generate less than 68 000 of steam per hour, whereas the watertube boilers tend to have a higher output. The overall efficiency of a boiler is related to the effectiveness of contacting water with the source of heat (i.e. coal, gas, oil, etc.).

Firatube boilers consist of a bundle of tubes set into a water drum. Water circulates around the outside of the tubes where it changes to steam and rises to the top of the drum and exits through a steam header. In watertube boilers, water circulates inside the tubes and the flame and hot combustion gases flow across the outside of the tubes. Steam forms on the heated side of the tubes producing a steam–water mixture that is less dense than the water on the unheated sides. Water circulation occurs either through density differentials or is controlled using a pump. The use of mechanical circulation provides more consistent control, particularly at high pressures where the density of steam and water are similar (Kemmer, 1988).

3.1.4 Sources of water used for cooling operations and boiler feedwater

Water used for cooling operations can come from a variety of sources including surface water (streams, lakes, or impoundments), groundwater, seawater, estuarine water, or reclaimed water. In selecting a water source, the major requirement is that an adequate supply of water is available to accomplish cooling. The water quality of the source dictates the need for pre-treatment and the feasibility of recycling the water with or without sidestream treatment. Water quality also influences the economic feasibility of using the water source for boiler feedwater or closed recirculating systems (Baron et al., 2000; Wijesinghe et al., 1996).

Because of the wide range of water sources that can be employed for cooling operations, water quality guidelines have been developed (Crook et al., 1994). A summary of these guidelines is given in Table 3.2 for cooling and boiler water. Water quality guidelines are based on prevention or minimisation of corrosion, scale or fouling. Boiler water quality requirements are more stringent than those for cooling water due to the relatively higher temperature and pressure conditions needed for steam production. In general, boilers that operate under higher temperature and pressure conditions require higher quality feedwater, particularly if the heat source is in direct contact with the tube, as in watertube designs. To prevent operational problems it is important to control the salt content of boiler water, as generally indicated by the conductivity, which then generally
Table 3.2  Water quality guidelines for cooling and boiler feedwater, and water quality associated with estuarine and reclaimed water

<table>
<thead>
<tr>
<th>Water quality parameter (all as mg l⁻¹ unless otherwise stated)</th>
<th>Cooling waterᵃ</th>
<th>Boiler e feedwaterᵇ,ᵇ</th>
<th>Estuarine waterᶜ</th>
<th>Reclaimed waterᵈ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional parameters</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alkalinity, mg l⁻¹ as CaCO₃</td>
<td>350</td>
<td>40–350</td>
<td>880–140</td>
<td>100–400</td>
</tr>
<tr>
<td>Hardness, mg l⁻¹ as CaCO₃</td>
<td>600</td>
<td>0.07–350</td>
<td>4300–5800</td>
<td>90–150</td>
</tr>
<tr>
<td>pH</td>
<td>6.9–9.0</td>
<td>7–10</td>
<td>7.5–8.6</td>
<td>6.6–7.5</td>
</tr>
<tr>
<td>Total dissolved solids</td>
<td>500</td>
<td>200–700</td>
<td>19000–24000</td>
<td>400–950</td>
</tr>
<tr>
<td>TSS</td>
<td>100</td>
<td>0.5–10</td>
<td>1–50</td>
<td>1–14</td>
</tr>
<tr>
<td>Turbidity, NTU</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organics</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C-BOD₅ᵉ</td>
<td>25</td>
<td>1–50</td>
<td>1–30</td>
<td></td>
</tr>
<tr>
<td>CODᶠ</td>
<td>75</td>
<td>1–5</td>
<td>5–100</td>
<td>20–80</td>
</tr>
<tr>
<td>Nutrients</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ammonia, mg l⁻¹ as N</td>
<td>1</td>
<td>0.1</td>
<td>0.1–5</td>
<td>0.2–50</td>
</tr>
<tr>
<td>Phosphate, mg l⁻¹ as P</td>
<td>4</td>
<td>0.1–5</td>
<td></td>
<td>2–60</td>
</tr>
<tr>
<td>Dissolved minerals</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aluminium</td>
<td>0.1</td>
<td>0.01–5</td>
<td>0.008–0.65</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Bicarbonate</td>
<td>24</td>
<td>48–170</td>
<td>110–160</td>
<td>20–200</td>
</tr>
<tr>
<td>Chloride</td>
<td>500</td>
<td>Variable</td>
<td>13000–18000</td>
<td>125–173</td>
</tr>
<tr>
<td>Iron</td>
<td>0.5</td>
<td>0.05–1</td>
<td>0.001–0.009</td>
<td>0.9–3</td>
</tr>
<tr>
<td>Manganese</td>
<td>0.5</td>
<td>0.01–0.3</td>
<td>&lt;1</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Silica</td>
<td>50</td>
<td>0.7–30</td>
<td>0.2–2</td>
<td>9–34</td>
</tr>
<tr>
<td>Sulphate</td>
<td>200</td>
<td>Variable</td>
<td>1700–2300</td>
<td>50–500</td>
</tr>
</tbody>
</table>

ᵃ Adapted from Crook et al. (1994).
ᵇ Range is based on boiler operating pressures.
ᶜ Water quality representative of wastewater reclamation facilities producing water for irrigation and cooling.
ᵈ Water quality representative of Tampa Bay, Florida.
ᵉ Five-day carbonaceous biochemical oxygen demand.
ᶠ Chemical oxygen demand.

determines the operational pressure (Fig. 3.3). Under low pressures, boilers can tolerate conductivity levels over 5000 µS cm⁻¹. However, as the operating pressure increases, the salt concentration becomes more critical. Thus, system design is based on the composition of the water available for use as boiler water.

Other critical water quality components for boiler water include alkalinity, silica, iron, manganese and copper (Fig. 3.4). Two operational problems are associated with high levels of alkalinity. As the water temperature increases, carbon dioxide is released increasing the potential for corrosion. In addition, carbonates can contribute to foaming that leads to deposits in the superheater, reheater and/or turbines (Puckorius, 1997; Troscinski and Watson, 1970; Vanderpool, 2001).

### 3.1.5 Water quality comparisons

For the purpose of comparison, water quality characteristics of estuarine water and reclaimed water are included in Table 3.2. Water quality associated with
freshwater sources, such as surface water or groundwater, is dependent on the local geohydrology, but would normally be similar to reclaimed water. As shown, there are differences in the content of nutrients, organics, and salts among the water sources. Seawater has higher levels of dissolved minerals than estuarine water.

The suitability of a water source for use in recirculating water systems as compared to once-through non-contact cooling is related to water quality and water availability. High levels of dissolved minerals and/or can demand additional treatment to prevent scale formation. For example, cooling systems that rely on estuarine or seawater tend to be non-recirculating, unless it is economical to provide on-site treatment for control of dissolved solids.
Typically, when reclaimed water is considered as a water source for cooling water, alternative sources such as surface water, groundwater, estuarine, or seawater may also be available locally. Thus the incentives for using reclaimed water are specific to the situation. When reclaimed water is used in lieu of surface or groundwater, the higher quality water sources can be preserved for other applications, such as drinking water. On the other hand, when reclaimed water is used in lieu of estuarine or seawater the impetus is usually related to discharge limitations.

Depending on the water source, water quality can vary seasonally. Groundwater tends to have fairly consistent characteristics, whereas the quality of surface water, estuarine water, seawater, and reclaimed water can be influenced by patterns of rainfall, runoff and evaporation, leading to significant seasonal variations in oxygen demand and suspended material (Fig. 3.5), nutrient levels (Fig. 3.6) and chloride (Fig. 3.7). Although water quality is source specific, the variation in wastewater characteristics and treatment alternatives means that similar trends arise with all reclaimed water sources.

Levels of dissolved minerals associated with estuarine water are shown for sodium and chloride in Fig. 3.8 and for calcium, magnesium, potassium, and sulphate in Fig. 3.9. Dissolved solids levels in estuarine waters are almost two orders of magnitude higher and sulphate and magnesium levels one order of magnitude higher than those levels associated with freshwater or reclaimed water. Typically, the mineral content of seawater can be two- to three-fold higher than that associated with estuarine waters. These water quality characteristics influence the extent of treatment required to allow for use of recirculating systems.

Another characteristic of reclaimed water that is different from fresh or saline water sources is the potential presence of a disinfectant residual. Reclaimed water is treated to meet requirements pertaining to microbiological safety. As such, disinfection is a key component of the treatment system. When chlorine is used for this duty, residual chlorine is usually present in the reclaimed

![Graph](image.png)

*Figure 3.5 Seasonal variations in five-day carbonaceous biochemical oxygen demand (C-BOD₅) and suspended solids (TSS) for a reclaimed water (data from St. Petersburg, Florida)*
Membranes for Industrial Wastewater Recovery and Re-use

Figure 3.6 Seasonal variations in ammonia (NH$_4$-N) and orthophosphate (PO$_4$-P) for a reclaimed water without nutrient removal (data from St. Petersburg, Florida)

Figure 3.7 Seasonal variations in chloride levels for a reclaimed water (data from St. Petersburg, Florida)

water. For cooling water applications, the presence of this residual disinfectant can act as a biocide and help to prevent biological fouling of the cooling system. There is also little seasonal variation in total residual chlorine (Fig. 3.10), provided chlorine is automatically dosed on demand through feedback control. Chlorine levels associated with reclaimed water thus depend on the operating practices of a given facility and, therefore, may differ from the trends shown in Fig. 3.10. In cases where disinfection is accomplished using ultraviolet (UV) irradiation or ozonation, residual disinfectants will not be present in the reclaimed water (Levine et al., 2002).
### 3.1.6 Optimisation of water use in recirculating cooling systems

Optimisation of water use in recirculating cooling towers is based on the quality of water entering and leaving the system. As water evaporates, dissolved constituents and salts become more concentrated in the liquid stream. The water quality of the recirculating stream must be controlled to prevent operational problems such as development of deposits on heat exchanger surfaces (scaling), corrosion, or biological fouling. To control the quality of the recirculating stream, water is removed as blowdown water, and to compensate for loss of water through blowdown, evaporation and drift water is added to the recirculating stream as make-up water (Table 3.3). Drift occurs when the water droplets become entrained in the discharge air stream; evaporation is from air passing through the cooling water and absorbing heat and mass; blowdown is the imposed bleed-off of water to reduce the concentration of contaminants. Continuous blowdown is the continuous removal of water, whereas intermittent blowdown is initiated manually or by feedback based on water quality. These same concepts apply to management of water quality for boiler systems (Asano et al., 1988, Burger, 1979; Kemmer, 1988; Puckorius and Hess, 1991).

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**Figure 3.8** Comparison of chloride and sodium levels in estuarine water (data from Tampa Bay, Florida)

**Figure 3.9** Comparison of calcium, magnesium, potassium, and sulphate levels in estuarine water (data from Tampa Bay, Florida)
To operate recirculating systems efficiently, it is important to prevent deposition (scaling) or fouling within the tower or heat exchangers. Water quality characteristics relating to mineral precipitation include calcium, magnesium, sulphate, phosphate, silica, pH, and alkalinity. As water evaporates, the concentration of dissolved constituents increases to the point where the solubility limit of mineral precipitates is exceeded within the recirculating water, particularly for carbonate scales at elevated temperatures. The solubility of mineral precipitates can be controlled by manipulating the pH of the recirculating water, addition of scale-control chemicals, and/or replacement of a portion of the recirculating water with less concentrated water. The considerations are thus identical to those of the operation of reverse osmosis plant (Section 2.4.3).

The quantity of water that must be removed as blowdown water can be calculated from a mass balance. The ratio of the concentration of a constituent in water to its concentration in the make-up water is called the cycles of concentration or concentration ratio, $R_C$, where:

$$ R_C = \frac{C_b}{C_m} $$

From an operations and water conservation perspective, it is desirable to have as high a $R_C$ value as possible. In general, the optimum $R_C$ is based on the chemical composition of the water and the solubility of the dissolved minerals. In some cases the $R_C$ is limited by calcium precipitation, such as calcium sulphate or phosphate. In other cases $R_C$ is limited by silica, magnesium, or other minerals. Thus, the characteristics of the make-up or source water can influence the maximum feasible $R_C$. Once the $R_C$ is determined, it can be used to determine the required make-up flow:
### Table 3.3 Summary of terms used to define flows through recirculating cooling towers

<table>
<thead>
<tr>
<th>Flow component</th>
<th>Definition</th>
<th>Flow rate, m(^3) h(^{-1})</th>
<th>Concentration of dissolved constituent(s), mg l(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recirculating water</td>
<td>Flow rate of water that is recirculating in tower</td>
<td>Q(_R)</td>
<td>CR</td>
</tr>
<tr>
<td>Blowdown</td>
<td>Flow rate of water deliberately removed from system to limit salt build-up</td>
<td>Q(_b)</td>
<td>C(_b) = C(_R)</td>
</tr>
<tr>
<td>Evaporation</td>
<td>Water lost to atmosphere due to evaporation: (\sim 2%) of recirculating water flow per 10°C temperature drop</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Q_e = Q_r \Delta T/700)</td>
<td>C(_e) = 0</td>
</tr>
<tr>
<td>Drift</td>
<td>Water lost due to entrainment in wind. When salt or brackish water is used, drift may contain high concentrations of salts and/or minerals that can deposit on crops, soils, or structures</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Q_d \sim 0.2%) for old units; (Q_d &lt; 0.005%) for efficient designs</td>
<td></td>
</tr>
<tr>
<td>Make-up water</td>
<td>Flow rate of water added to flow stream to replace water by evaporation, drift and blowdown</td>
<td>(Q_m = Q_e + Q_d + Q_b + Q_t)</td>
<td>C(<em>m) = C(</em>{source\ water})</td>
</tr>
<tr>
<td>Other losses</td>
<td>Losses of water due to leaks</td>
<td>Q(_l)</td>
<td>C(_l) = C(_R)</td>
</tr>
<tr>
<td>System capacity</td>
<td>Total volume of water in system</td>
<td>V</td>
<td></td>
</tr>
<tr>
<td>Temperature differential</td>
<td>Difference between average water temperature following evaporation and average water temperature returning to tower</td>
<td>(\Delta T = T_{inflow} - T_{return}, ^\circ C)</td>
<td></td>
</tr>
<tr>
<td>Cycle time</td>
<td>Time required for water to travel around circulating loop</td>
<td>(t_c = V/Q_R)</td>
<td></td>
</tr>
<tr>
<td>Holding time index</td>
<td>The half-life of chemicals added to the system</td>
<td>(t_{1/2} = (\ln 2) \times V \times (R_c \cdot 1/E))</td>
<td></td>
</tr>
</tbody>
</table>
The extent of recirculation can be increased by using chemical treatments either to adjust the pH of the water or to sequester minerals and prevent deposition, in the same way as reagents are used to ameliorate scaling in reverse osmosis plant (Table 2.15). In most cases, the $R_c$ is optimised based on water availability, water quality, and treatment costs. Typically, evaporative recirculating cooling towers that use reclaimed water operate with $R_c$ values ranging from 1 to 3. An advantage of higher $R_c$ values is that reduced quantities of make-up water are required and there is less blowdown water to be treated or discharged. It should be noted that the concentration of dissolved constituents in the blowdown water increases with increasing $R_c$. Therefore, one trade-off associated with higher $R_c$ levels is the increasing costs of disposal or treatment of the blowdown water.

The impact of recirculation on water quality in a recirculating cooling system using reclaimed water as a source water is shown in Fig. 3.11. This system is operated with a $R_c$ of about 2. A comparison of the conductivity of the source water to that of the recirculating stream reveals that the concentration of dissolved solids increases about two-fold in this system. When blowdown water is removed from this system it is returned to the wastewater reclamation facility. Higher $R_c$ values for this source would yield a blowdown water with a higher salt content that might preclude discharge to the wastewater reclamation facility.

### 3.1.7 Cooling tower water quality issues

Industrial cooling tower operations are susceptible to four potential water quality problems: (1) scaling, (2) biological growth, (3) fouling of the heat transfer surfaces, and (4) corrosion. Each of these issues can affect the performance and efficiency of the cooling system. Scaling can reduce the heat transfer rate by blocking the pipe walls and reducing the water flow. Biological growth can further reduce the heat transfer rate and can also lead to the formation of biofilms that can clog the system. Fouling of the heat transfer surfaces can also reduce the heat transfer rate by blocking the flow of water and can increase the pressure drop in the system. Corrosion can damage the system and can also lead to the release of metals into the water, which can be toxic to the environment.

$$Q_m = R_c \times Q_b$$  \hspace{1cm} (3.2)
exchangers and condensers and (4) metallic corrosion. These water quality problems can result from any water source (fresh, reclaimed, or salt) unless appropriate preventive measures are incorporated into the cooling water system. Definitions of each cooling water quality issue are given below. Chemicals such as chlorine and chelating agents are added to prevent biofouling and inhibit mineral build-up. As the water volume is reduced through evaporation and drift, the concentration of these chemicals and their by-products increases. Cooling towers can also contain chemicals from the ambient air.

Scaling
Scaling refers to the formation of mineral deposits, usually on hot surfaces, which can compromise heat exchanger efficiency. As with dense membrane processes, calcium deposits (calcium carbonate, calcium sulphate and calcium phosphate) are the predominant form of scale. Such deposits are somewhat heterogeneous in nature (Fig. 3.12) and their accumulation on the surfaces of heat exchangers can reduce the heat transfer efficiency, causing overheating of metal and, ultimately, boiler tube failure. In addition, the presence of deposits can provide habitats for the growth of microorganisms within the cooling system. Magnesium scales (magnesium carbonate and phosphate) can cause a similar problem. Silica scales are particularly problematic, since silica is largely insoluble and forms very tenacious deposits. Silica can volatilise at the high temperatures of boiler systems and become entrained in the steam (Dyson, 2001; Troscinski and Watson, 1970; Vanderpool, 2001). As the steam cools within the turbine, the silica can crystallise on the turbine nozzles or blades leading to increased frictional resistance and reduced steam velocities. Because the deposits are not uniformly distributed, the turbine rotors become imbalanced and produce excessive vibrations.

![Figure 3.12](image)  
*Figure 3.12  Electron micrograph of calcium deposit from cooling tower operations. The length of the white bar represents 10 µm*
Scale composition depends on the relative mineral content of the recirculating stream. A summary of solubility constants associated with common mineral precipitates has been presented in Table 2.14. While the chemical content of each system has been a function of the source water and the $R_c$, it is possible to apply chemical equilibrium models, such as Argo Analyser (Fig. 4.7) to evaluate the potential for scale formation. As a first approximation, the solubility relationships can be used to identify the chemical constituents that are likely to form deposits. Typically, when reclaimed water is used as a source water, the first calcium salt to precipitate is calcium phosphate unless the water has been pre-treated for phosphorus removal.

Scale control can be accomplished through chemical precipitation followed by solids removal (sedimentation, filtration, etc.) to reduce the concentration of minerals in the recirculating stream. Chemicals used to promote upstream precipitation include lime, caustic soda, alum, and various formulations of organic or inorganic polymers. Acidification or addition of scale inhibitors can control scaling by increasing the solubility of minerals in the recirculating stream. Phosphate specifically can be removed biologically, and multivalent ions may be removed by ion exchange.

The solubility of mineral precipitates that form from hydroxides, phosphates, or carbonates typically increases with decreasing pH. To prevent scale formation, the pH of the water is reduced to about 7 using sulphuric acid. The additional sulphate and lower pH convert calcium and magnesium carbonates into more soluble sulphate compounds. It is important to control the amount of acid added to maintain some residual alkalinity in the system, since excess acid can cause accelerated corrosion. Acids used to control pH of the recirculating stream include sulphuric, hydrochloric, and citric acids. Alternatively gases can be used to acidify the water such as carbon or sulphur dioxide. Chemical chelators such as ethylenediamine tetraacetic acid (EDTA) and polymeric inorganic phosphates can also be added, often in-line (Fig. 3.13), to increase the solubility of scale forming constituents.

**Biological growth**

The warm, moist environment in cooling towers coupled with the availability of nitrogen, phosphorus, and organics provides an ideal environment for microbial growth. Typically, microbial growth results in biofilm formation and fouling, in which microbial products encourage the attachment and growth of heterogeneous deposits containing both microorganisms and inert materials, on heat exchanger surfaces. These biofilms then interfere with heat transfer and water flow. During extended operating periods, portions of the biofilm slough off of the surface. This microbial biomass contains particles and other debris that can settle, further inhibiting effective heat transfer. Some types of microorganisms release corrosive by-products during their growth such as organic acids (e.g. acetic) or inorganic acids (e.g. hydrogen sulphide) leading to microbially induced corrosion (MIC), a phenomenon exacerbated by standing water conditions.

Bacteria that may be present in cooling water include *Pseudomonas, Klebsiella,Enterobacter, Acinetobacter, Bacillus, Aeromonas, and Legionella* (Adams et al.,
1978; Wiatr, 2002). Once a biofilm forms, it provides a protective habitat for microorganisms (Fig. 3.14). Biocides can be used to control biofilms as part of the internal chemical treatment process, the type and required dosage depending on the organic and nutrient content of the make-up water. The most commonly used biocide is chlorine, though other chemical approaches are also effective. Ozone is a powerful biocide effective for control of bacteria, viruses, and protozoa, but can exacerbate problems of scale adhesion since by-products from the oxidation of biofilms can serve as binding agents for scale on heat exchanger surfaces.

When reclaimed water is used for cooling, the assurance of adequate disinfection is a primary concern to protect the health of workers and individuals exposed to aerosols from the cooling towers. The disinfection requirements for the use of reclaimed water in cooling towers are site specific and based on the potential for exposure to aerosols from cooling operations and prevention of biofilm growth. Limited data are available on relative quantities of microorganisms in recirculating cooling systems. Pathogen survival depends on the source water quality, pretreatment mechanisms, and the type and dosages of biocides used in the facility (Levine et al., 2002). While there are no universal standards, the most frequently monitored bacteria include total and faecal coliforms and Legionella pneumophila. Typically disease outbreaks are associated with levels over 1000 cfu (colony forming units) per ml in cooling towers. A comparison of the levels of Legionella pneumophila in recirculated cooling water is shown in Fig. 3.15. This facility uses a pro-active approach by conducting quarterly monitoring. Typical values range from non-detectable to 300 cfu ml⁻¹. Monitoring can provide insight into the effectiveness of disinfection practices.

Water velocities below 0.3 m s⁻¹ and temperatures over 50°C tend to promote biofilm formation and the associated fouling reactions. Control of fouling is
achieved by the addition of chemical dispersants to prevent particles from aggregating and subsequently settling. Also, a secondary benefit of chemical coagulation and filtration processes is the removal of some contaminants that contribute to fouling.

The critical part played by biofouling in determining wastewater reuse capability has been recognised in extensive studies carried out in Grangemouth, Scotland (Glen, 2002). Grangemouth is one of the largest industrial complexes in the UK, with major petrochemical industries concentrated in a small area. Water consumption is 26 million m³ per year, costing business around $24 m per annum at (2002 dollars), with one of the main uses being for heat exchange
systems. To assess fouling propensity of possible feedwater sources the National Engineering Laboratory based at Grangemouth have developed device for assessing biofilm formation through thermal resistance measurements. Results so far have revealed fouling rates of recovered municipal effluent to be generally higher than other sources, such as canal water, and highly dependent on flow velocity. On the other hand, examples exist of employing membrane treatment for the production of boiler feedwater from secondary municipal effluent (Section 5.1).

**Metallic corrosion**

In cooling systems, corrosion can occur when an electrical potential between dissimilar metal surfaces is created. The corrosion cell consists of an anode, where oxidation of one metal occurs, and a cathode, where reduction of another metal takes place. The presence of dissolved minerals can accelerate the corrosion reaction. Dissolved oxygen and certain metals (manganese, iron, and aluminium), which can arise in cooling intakes depending on the water source (Salvito et al., 2001), promote corrosion because of their relatively high oxidation potentials.

Mild steel and copper alloys are used extensively in heat exchanger systems due to their excellent heat transfer capabilities. However, the build-up of minerals in cooling water tends to be aggressive towards these materials. Analysis of corrosion is complex due to the interplay of water quality variables that can either induce or prevent corrosion. A summary of water quality determinants significant for corrosion control is given in Table 3.4. The corrosion potential of cooling water can be controlled by the addition of chemical corrosion inhibitors. The concentration of chemicals in the recirculating stream, increases with increasing $R_C$ and may require removal from blowdown water prior to discharge.

### 3.1.8 Governing legislation and guidelines

Regulatory requirements and industrial guidelines relating to the use of water for cooling and boiler applications are fairly localised. In the USA regulatory requirements address water intake structures, air emissions, and discharge of blowdown water. Currently, under Section 316(b) of the Clean Water Act, it has been proposed to implement more stringent protection measures at water intake structures to protect fish, shellfish, and other aquatic life (USEPA, 2002a). This proposed regulation could lead to the need for retrofitting of once-through cooling water intakes and ultimately lead to increased use of recirculating systems.

Airborne emissions from coal-fired power plants and waste-to-energy facilities are a subject of increasing scrutiny in urban environments. Increased emissions control requirements coupled with uncertainties about the long-term availability of fossil fuels may result in changes in the distribution of energy sources. As coal-fired plants are displaced by alternative fuels, changes will also occur in the quantities of cooling water required. It has been estimated that conversion from coal-fired power plants to alternative fuels could reduce evaporative consumption by 25% (Powicki, 2002).
<table>
<thead>
<tr>
<th>Parameter, optimum range</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkalinity, consistency</td>
<td>Variations in concentration can influence formation of metal carbonate complexes and result in increased metal release.</td>
</tr>
<tr>
<td>Ammonia, 0 mg L(^{-1})</td>
<td>Can complex with metals, increasing solubility and metal release, and can also promote biological growth and biocorrosion. Corrosive to copper alloys causing metal loss and stress corrosion cracking at levels greater than 2 mg L(^{-1}) as N. Reduces effectiveness of chlorine through chloramine formation. Increases with cycles of concentration.</td>
</tr>
<tr>
<td>Calcium, consistency</td>
<td>Can form calcite on pipe walls, possibly preventing corrosion. Variations in water quality promotes dissolution of scales and thus metal release.</td>
</tr>
<tr>
<td>Chloride, consistency</td>
<td>Can increase corrosion rates under conditions of low dissolved oxygen. Can form complexes with metals increasing solubility and metal release. Influences corrosion of most metals including mild steel, copper, stainless steel. Chlorides over 200 mg L(^{-1}) can cause stress corrosion cracking and/or pitting corrosion.</td>
</tr>
<tr>
<td>Dissolved oxygen, consistency</td>
<td>Can serve as an electron acceptor for corrosion reactions. However it also reacts to form protective oxide layers that prevent corrosion. Localized differences can promote corrosion.</td>
</tr>
<tr>
<td>Iron, consistency</td>
<td>Can combine with phosphate and can also deactivate polymers used to inhibit calcium phosphate scaling.</td>
</tr>
<tr>
<td>Magnesium, consistency</td>
<td>Can interfere with deposition of calcium complexes on pipe wall.</td>
</tr>
<tr>
<td>Organic carbon (TOC), consistency</td>
<td>Can coat pipe surfaces and prevent metal release. It can also form metal complexes and increase metal release and can promote biological growth and biocorrosion.</td>
</tr>
<tr>
<td>Orthophosphate, 0.5–5 mg L(^{-1}) as PO(_4)</td>
<td>Can form a passivating layer on the internal surface of heat exchangers.</td>
</tr>
<tr>
<td>pH, 7.3–7.8 pH units</td>
<td>Acidity increases metal solubility and promotes CO(_2) liberation from alkalinity. pH may be locally higher at pipe surface due to OH(^-) generation.</td>
</tr>
<tr>
<td>Silica, below 150 ppm</td>
<td>Can act to passivate surfaces, depending on the concentration, thus decreasing the potential for metal release. Total silica refers to colloidal and ionic silica in water. Polymeric silica is non-reactive but it can revert to silica monomers after heating in boiler water.</td>
</tr>
<tr>
<td>Sulphate, consistency</td>
<td>Can interfere with formation of some scales, thus increasing the potential for metal release.</td>
</tr>
<tr>
<td>Temperature, consistency</td>
<td>Corrosion rates increase with increasing temperature. Influences solubility, rates of microbiological activity, water density and associated mixing efficiency.</td>
</tr>
</tbody>
</table>

### 3.1.9 Volumes and quality of aqueous process waste streams

Aqueous process waste streams include blowdown water from cooling operations and boilers. The quantity of water generated is related to operational variables including the source of the make-up water and the cycles of
Concentration. Treatment requirements depend on discharge requirements or incentives for reuse of the water. Typically discharge standards are based on pH, temperature, C-BOD₅, TSS, toxic substances, and dissolved solids.

Several approaches are used for treatment of water associated with thermoelectric power generation depending on the source of the cooling and boiler water, the type of cooling system, and the discharge requirements. The characteristics of the water used for make-up dictate the need for supplemental treatment. Only minimal levels of treatment are normally required for once-through cooling systems. The main water quality objective is to prevent scaling, corrosion or fouling. Because of the relatively short contact time and relatively low temperatures, effective cooling can be accomplished using chemical amendments to act as corrosion and scale inhibitors. One of the factors limiting the use of chemical amendments for once-through systems is that the water is typically discharged back to the environment. Thus, the type of chemicals that are used must meet local water quality requirements for discharges (Piller and Coupe, 2001).

Recirculating systems are more vulnerable to scaling, corrosion and fouling due to accumulation of solutes. Thus, treatment objectives are targeted at reducing the mineral, nutrient, and/or organic content of the make-up water. Management practices for the blowdown water include discharge to a wastewater reclamation facility, blending with other waste streams, or treatment either to recover the water for reuse or to meet discharge requirements through the use of various biological and physicochemical processes (Table 3.5). Historically, the predominant treatment systems have been chemical-based treatments including lime softening and chemical precipitation using metal salts. Recently, the use of ion exchange, membrane systems, and thermal processes has become more cost-competitive. The new generation of treatment technologies, largely membrane based, have the advantage of producing a high-quality product water that can be reused for duties such as boiler feedwater (Sections 5.1 and 5.2). Disposal of brines and waste streams arising from these processes remains a challenge, however, and if zero liquid discharge is demanded recourse is inevitably made to evaporation, sometimes preceded by electrodialysis (Section 5.3).

### 3.1.10 Current reuse practices and opportunities

The extent to which reclaimed water is used for cooling water and boiler water applications depends on several factors. Firstly, the reclaimed water source must be within a practical distance from the industrial, ideally less than 50 km. Secondly, economic and environmental factors should be favourable for the use of reclaimed water. The key issues are the relative availability/scarcity of alternative sources of cooling water. In addition to the need for water, when the use of once-through cooling is limited by regulatory or discharge requirements reclaimed water can become a viable alternative. To date, the use of reclaimed water is fairly well accepted throughout the world, and some examples are summarised in Table 3.6.
<table>
<thead>
<tr>
<th>Treatment goal(s)</th>
<th>Treatment approach</th>
<th>Description</th>
<th>Treatment application</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reduce scale-forming potential, remove</td>
<td>Lime softening</td>
<td>Use of lime to precipitate cations, metals, and phosphorus from solution</td>
<td>X X X</td>
<td>Chloride and sulphate ion pairs can limit precipitation of calcium carbonate, Sludge requires treatment/disposal</td>
</tr>
<tr>
<td>phosphorus, increase pH</td>
<td></td>
<td></td>
<td></td>
<td>Need source of heat. Sludge requires treatment/disposal</td>
</tr>
<tr>
<td>Removal/conversion of nitrogen; removal</td>
<td>Hot process softening</td>
<td>Use of lime and soda ash under elevated temperature conditions</td>
<td>X X</td>
<td>Not effective in high salt conditions</td>
</tr>
<tr>
<td>of phosphorus</td>
<td>Biological nutrient removal</td>
<td>Combination of aerobic, anoxic, and anaerobic processes to optimise conversion of organic and ammonia nitrogen to molecular nitrogen (N₂) and removal of phosphorus</td>
<td>X X</td>
<td></td>
</tr>
<tr>
<td>Removal of calcium, magnesium, iron,</td>
<td>Coagulation and precipitation</td>
<td>Use of aluminium or iron salts to promote precipitation of phosphorus</td>
<td>X</td>
<td>Sludge requires disposal/treatment</td>
</tr>
<tr>
<td>ammonium, nitrate, and other dissolved</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ions</td>
<td>Ion exchange</td>
<td>Exchange of ions between an exchange resin and water using a flow-through reactor</td>
<td>X X X</td>
<td>Regenerant requires disposal</td>
</tr>
<tr>
<td>Electrodialysis reversal (Section 2.4.2)</td>
<td></td>
<td>Membrane-based process using an electric field to separate ionic constituents across alternating anion-selective and cation-semi-permeable membranes</td>
<td>X X</td>
<td>Requires conductivity &gt; 4000 mS cm⁻¹. Not effective for non-ionised species such as silica, ammonia, and carbon dioxide</td>
</tr>
<tr>
<td>Treatment goal(s)</td>
<td>Treatment approach</td>
<td>Description</td>
<td>Treatment application</td>
<td>Limitations</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>-----------------------------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Removal of dissolved solids, organics, pathogens</td>
<td>Reverse osmosis</td>
<td>Membrane system to separate ions from solution based on reversing osmotic pressure differentials</td>
<td>X</td>
<td>Potential for fouling of RO membranes. Brine requires treatment/disposal</td>
</tr>
<tr>
<td></td>
<td>Nanofiltration and ultrafiltration</td>
<td>Pressure-driven membrane processes to remove constituents based on charge and molecular size</td>
<td>X</td>
<td>Potential for fouling of membranes. Reject stream requires treatment/disposal</td>
</tr>
<tr>
<td>Facility (Reference)</td>
<td>Application</td>
<td>Method</td>
<td>Water (Power) offset</td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
<td>-------------</td>
<td>--------</td>
<td>---------------------</td>
<td></td>
</tr>
<tr>
<td>Palo Verde nuclear power plant, Arizona (Blackson and Moreland, 1998)</td>
<td>Cooling water for 3800 MW nuclear power plant using nitrified reclaimed water with supplemental chemical treatment: softening, phosphorus removal, pH adjustment, and chlorination</td>
<td>Recirculating cooling towers with 18–23 cycles of concentration; zero liquid discharge; evaporation ponds for blowdown water</td>
<td>&gt; 250 000 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Chevron Oil Refinery, Oakland, California (Puckorius, 1997)</td>
<td>Refinery cooling water using reclaimed water with supplemental lime treatment, filtration, and disinfection</td>
<td>Recirculating cooling tower</td>
<td>17 000 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Lockheed-Martin Electronics operation, Ocala, Florida</td>
<td>Cooling water for production of electronics assemblies and secondary cooling of HVAC using manufacturing process wastewater treated with hydrogen peroxide and ozone</td>
<td>Counter current cooling towers</td>
<td>6800 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Bergen generating facility, Bergen County, New Jersey (Puckorius, 1997)</td>
<td>Cooling water for 625 MW thermoelectric power plant using reclaimed water treated with biocide, acid, and polymer</td>
<td>Recirculation cooling; blowdown discharged to sanitary sewer</td>
<td>5400 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>City of Tampa waste-to-energy facility, Tampa, Florida</td>
<td>22 MW facility. Processes 1000 tons per day</td>
<td>Two cell recirculating evaporative cooling tower. Blowdown water is discharged to wastewater treatment plant</td>
<td>2200 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Oil refinery, Singapore</td>
<td>Oil refinery using reclaimed water treated with acid, scale inhibitor, dispersant, and corrosion inhibitors</td>
<td>Once-through cooling of 32 water-cooled pumps and two recirculating cooling towers</td>
<td>1600 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Flag Fen, UK</td>
<td>Dual membrane MF: 28 modules, 0.1 µm pore size; RO: 4:2 array; 80% recovery</td>
<td>Boiler feedwater using polished municipal</td>
<td>1200 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Facility (Reference)</td>
<td>Application</td>
<td>Method</td>
<td>Water (Power) offset</td>
<td></td>
</tr>
<tr>
<td>-----------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------</td>
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</tr>
<tr>
<td>Guadelope power plant, Guadelope County, Texas (Prato et al., 2001)</td>
<td>Natural gas, 1000 MW. Make-up water treated by softening; blowdown treated by evaporation, crystallization and EDI to produce boiler feedwater</td>
<td>Recirculating cooling tower</td>
<td>820 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>Raytheon Corporation, St. Petersburg, Florida</td>
<td>Air conditioning cooling towers</td>
<td>Three recirculating cooling towers</td>
<td>500 m³ d⁻¹</td>
<td></td>
</tr>
<tr>
<td>PECO Energy nuclear generating station, Limerick, Pennsylvania (PECO, 2002)</td>
<td>Cooling water for 1200 MW nuclear power plant</td>
<td>Recirculation cooling system with 3–6 cycles of concentrations</td>
<td>(0.85 m³ MWh⁻¹)</td>
<td></td>
</tr>
<tr>
<td>International Microelectronics, North San Jose, California (USEPA, 2002b)</td>
<td>Water used to cool equipment for microelectronics manufacture</td>
<td>Closed loop cooling system with circulating chilled</td>
<td>18–28 m³ d⁻¹</td>
<td></td>
</tr>
</tbody>
</table>
It is evident that as membrane technologies advance and the efficiency of removing salts and other impurities from water increases, the use of reclaimed water for cooling and boiler applications will increase. A promising opportunity is the development of cost-effective treatment systems for treatment of blowdown water from cooling and boiler applications to produce high-quality water for make-up water or other applications. Three case studies relating to power plants are detailed in Sections 5.1–5.3.

References


USEPA (2002a). Cooling water intake structures at large existing power plants. EPA-821-F-02-003.


3.2 The pulp and paper industry

3.2.1 Introduction

Pulp and paper production in the world is localised in countries with natural forest resources (Figs. 3.16 and 3.17). Thus most paper products are manufactured in the USA, Canada, China, Europe, Korea and Brazil. In spite of the growing popularity of the internet and electronic journals, global pulp and paper production is steadily increasing at the same rate as gross domestic product in the world. In 1998 the world production of pulp and paper was estimated to exceed 300 million tonnes for the first time.

The pulp used for paper products is either chemical or mechanical pulp, or recovered paper. The paper products, in which the different pulps are used in varying proportions, can be divided in the following paper grade groups (Zippel, 2001):

- Newsprint
- Fine paper ("wood-free paper")
- Magazine paper ("wood-containing paper")
- Tissue
- Packaging paper (liner and fluting)
- Multiply board

In addition to different pulps, different fillers and/or coating materials, such as pigments, are used. A survey of 100 German paper mills showed that the raw materials in papermaking were (Zippel, 2001):

- Chemical pulp – 21%
- Mechanical pulp – 14%
- Recovered paper – 43%
- Filler, pigments – 16%
- Water – 6%

The proportions, however, vary depending on the location of the paper mill and on the paper grade produced.

Even though water constitutes only 6% of the raw materials used in paper making, its importance is significantly enhanced by its role as a carrier of raw materials through the paper making process before the paper machine drying section. The water quality has a direct effect on process operability and product quality.

The specific freshwater consumption for the pulp and paper industry is defined as the amount of freshwater used for the manufacturing process, taken from external sources, related to the air-dry net weight material produced as finished or intermediate rolls. During the twentieth century the specific fresh water consumption has fallen dramatically from around 300 m$^3$/t (i.e. m$^3$ per tonne) at
the turn of the century to between 20 and 30 m³/t today. Naturally, an important factor affecting the water consumption is the paper grade as well as the technical age of the paper machine: the more modern the paper machine, the lower the freshwater consumption.

Traditionally flotation and evaporation have been widely used as separation and purification techniques in the pulp and paper industry. Since the 1980s membrane filtration has also been adapted to these applications. Moreover, along with increasing knowledge and better membranes and modules, membrane filtration has gradually become a more attractive option through its versatility, intensity (low footprint), and significantly lower energy consumption compared to evaporation.
3.2.2 Pulping and paper manufacturing processes

Wood is a renewable resource that consists mainly of cellulose (45%), hemicellulose (30%) and lignin (25%). Lignin is the resinous material that binds the cellulose fibres together. In addition, wood has less than 5% of substances like terpenes, resins and fatty oils and acids. Wood for papermaking has to be debarked using a large, rotating drum that agitates the logs and allows the bark to be stripped by the abrasive action of the neighbouring logs. The logs are then chipped using a large rotating blade prior to pulping.

The pulping process (Table 3.7) separates the cellulose and hemicellulose from the lignin and removes other tree oils and resins. The remaining fibres are used to produce the paper. There are two main pulping processes, chemical pulping and mechanical pulping. Chemical pulping produces very pure cellulose fibres, and is the most common form of pulping worldwide. In Kraft pulping, also called the sulphate process, the wood chips are heated in sodium hydroxide and sodium sulphide to produce a strong dark brown pulp. Because of the dark colour of the resulting pulp, Kraft fibres require considerable bleaching to make them usable. Kraft pulping is a highly efficient process for removing lignin and resins in softwoods while still producing a high-quality pulp. Wood wastes are burned for fuel and more than 95% of the pulping chemicals are recovered for reuse. In the sulphite process, less commonly used for pulping than the Kraft process but still widely used in Central Europe, the chips are heated in sulphuric acid. Recovery of the pulping chemicals is less well developed than for the Kraft process.

In mechanical pulping the debarked logs are forced against a grinding stone or rotating metal disks. As much as 95% of the wood resource is turned into pulp though this action, since the lignin and tree resins remain in the pulp (unlike Kraft pulping in which the yield is only 45–50%). The high lignin content in the pulp causes it to darken when exposed to sunlight. In mechanical pulping the chips can also be steam treated to soften them prior to grinding, a process known as thermomechanical pulping (TMP). In chemothermomechanical pulping (CTMP) the wood chips are soaked with sulphur-based chemicals prior to steaming to expedite the extraction of lignin and resin from the wood. Mechanical pulp is used for lower grade papers, such as newsprint and telephone directories, as well as for coated papers.

Bleaching is used to purify and clean the pulp. Bleaching removes lignin, which affects the purity of the fibre. Kraft mills use mainly chlorine dioxide or other strong oxidants, such as oxygen, ozone or hydrogen peroxide, to bleach pulp while most mechanical pulp bleaching operations employ hydrogen peroxide. Chlorine gas is still used in some old mills, particularly in USA, but due to its tendency to form polychlorinated organic compounds it has been mostly abandoned as a bleaching agent. Mechanical pulp is most often bleached with hydrogen peroxide or with thiosulphate.

Recovered paper is used as fibre source in papermaking, particularly in regions where the population density is very high. Worldwide, over 95 million tonnes of paper are recovered each year to be made into recycled paper and paperboard, contributing more than a third of the total fibre used to make the world's paper
### Table 3.7 Comparison of mechanical and chemical pulping processes (Kroesa, 1990)

<table>
<thead>
<tr>
<th>Process</th>
<th>Mechanical pulping</th>
<th>Chemical pulping</th>
</tr>
</thead>
<tbody>
<tr>
<td>Process</td>
<td>Groundwood log is pressed against a grindstone</td>
<td>CTMP&lt;sup&gt;a&lt;/sup&gt; wood chips are vapour-heated and chemically treated before grinding</td>
</tr>
<tr>
<td>Lignin</td>
<td>Much lignin left in pulp, turns dark when exposed to light</td>
<td>Little lignin left, remains white after bleaching</td>
</tr>
<tr>
<td>Fibre make-up</td>
<td>Yellow torn fibres, very weak</td>
<td>Slightly yellow, more whole fibres, fairly strong</td>
</tr>
<tr>
<td>Fibre make-up</td>
<td>Light brown whole fibres, strong, soft</td>
<td>Light brown whole fibres, strong, soft</td>
</tr>
<tr>
<td>Pulp yield</td>
<td>0.9–0.95 t pulp/t wood</td>
<td>0.45–0.5 t pulp/t wood</td>
</tr>
<tr>
<td>Bleaching</td>
<td>Hydrogen peroxide</td>
<td>Chlorine/chlorine dioxide, sometimes oxygen, hydrogen peroxide</td>
</tr>
<tr>
<td>Electrical energy</td>
<td>2000 kWh/t</td>
<td>1150 kWh/t</td>
</tr>
<tr>
<td>Steam energy</td>
<td>n/a</td>
<td>1000 kWh/tonne</td>
</tr>
<tr>
<td>Water</td>
<td>10 000–15 000 gallons/t</td>
<td>5200 kWh/t</td>
</tr>
<tr>
<td>Water</td>
<td>15 000–45 000 gallons/t</td>
<td>4400 kWh/t</td>
</tr>
</tbody>
</table>

<sup>a</sup> Chemothermomechanical/pulping.

(TAPPI, 2000). At the mill site the recovered, sorted paper is stored, and the paper is re-pulped, screened and cleaned. The printing ink and “stickies” (sticky materials like glue residue and adhesives) are removed in the de-inking process. After de-inking the pulp is refined and, if needed, bleached using hydrogen peroxide, chlorine dioxide, and/or oxygen.

Paper manufacturing involves three stages: stock preparation, pressing, and thermal drying. During stock preparation, wood fibre is mixed with a 1% solution of minerals in water and the water drained from the stock to form a web of paper. In the next step, pressing, the paper is squeezed between rollers and a felt (a large mat of nylon and polyester filaments) to produce a material generally containing 40–50% water. In the third step water is removed by thermal drying to produce paper containing 2–6% water. If size (a starch or starch-like substance used to modify the surface properties of the paper) is to be used a size press coating consisting of starch and minerals is put onto the paper next. Following this the paper is dried once again before finally being wound into a large roll. The surface of the paper is often modified by coating to improve the print quality: the paper is passed under a blade that deposits a homogeneous coating onto its surface. Different kinds of pigments and adhesives are used for different grades of paper. Supercalendering, used for some grades to make the paper glossy and smooth, employs a number of large rollers to simultaneously press and heat the paper. Sheet converting is the final stage of papermaking, in which the rolls of paper may be cut into individual sheets.

Pulp and paper mills can be either discrete production units or form an integral process. The water consumption differs between non-integrated and integrated...
mills. In a non-integrated paper mill, the purchased pulp is the major fibre source and is delivered at 90% solids. In an integrated paper mill, pulp from the pulp mill is stored at about 12–16 wt% solids. In both cases the web leaves the press section at about 40–46% solids content. Consequently, there is a net demand for mill water in a non-integrated paper mill and a net surplus of water in an integrated mill. However, exchange of water between the pulp and paper operations in an integrated mill can change the situation, even though such exchange is nowadays minimised. The production of mechanical pulp always exerts the greatest water demand.

3.2.3 Effluent volumes and quality

Over the years production techniques have been modified and made more efficient and cost effective. Thus the process itself, the paper grade and the technical age of the mill are all key factors when the effluent volumes and quality are considered. They also naturally depend very strongly on the pulping process and on the type of raw materials as well as on the paper grade to be produced (Table 3.8).

Pulping

In debarking significant amounts of bark, fibres and dissolved organic substances are released into the water. The debarking effluent is also toxic, since some of the compounds found in bark are water soluble and heavily toxic. Thus dry debarking or separate treatment is commonly practised, when possible.

In the Kraft chemical pulping process (Fig. 3.18) dissolved organic substances, mainly polysaccharides and lignin, are released in the cooking process. Organic lignin compounds with spent white liquor (the main components of which are Na₂S and NaOH, the pH level being around 14) form the black liquor that is removed from the washing stage. In the bleaching stage, the first alkaline extraction, E₁, extracts organic lignin compounds along with polysaccharides from the pulp. When chlorine-based bleaching agents, such as elementary

<table>
<thead>
<tr>
<th>Paper grade</th>
<th>Major raw materials used&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Newsprint</td>
<td>No added new fillers; few added chemicals; sometimes pigments added;</td>
</tr>
<tr>
<td></td>
<td>very little colour; uses mechanical pulp or RCF</td>
</tr>
<tr>
<td>Kraft wrapping</td>
<td>No added fillers; coloured; uses mainly unbleached kraft pulp</td>
</tr>
<tr>
<td>Kraftliner or testliner</td>
<td>No added fillers; uses unbleached kraft pulp and RCF or only RCF</td>
</tr>
<tr>
<td>Board</td>
<td>All fibre types including RCF</td>
</tr>
<tr>
<td>Writing and printing</td>
<td>All grades of fibre, mainly bleached; fillers; sizes; colours; brighteners; may be coated</td>
</tr>
<tr>
<td>Tissue</td>
<td>No added fillers; wet strength additives; uses chemical pulp + RCF in different mixtures</td>
</tr>
<tr>
<td>Speciality papers</td>
<td>The pulp used can vary considerably in type and quality</td>
</tr>
</tbody>
</table>

<sup>a</sup> RCF: recovered fibre.
chlorine or chlorine dioxide, are used some chlorinated substances are formed in bleaching. They are measured as AOX, i.e. adsorbable organic halogens. Some nutrients (phosphorus and nitrogen) as well as salts are also released during the bleaching operation.

An example of effluent stream quality for a modern softwood Kraft pulp mill is shown in Table 3.9 (Myrén, 1993). It is apparent from these data that the bleach plant is the major producer of effluent. However, even though the volume of the wood room effluent, i.e. from de-barking, is only about 5% of the total volume, it is nonetheless the most toxic effluent and constitutes 10% of the colour load in the total effluent. Moreover, conventional wastewater treatment by the activated sludge (AS) process removes only around half of the COD. AOX and phosphorus load from the effluent and leaves the colour substantially unremoved. From this it may be concluded that the colour is mostly formed by long-chain organics that are not well digested by the activated sludge plant. The specific characteristics of the effluents depend on the pulping process (Table 3.10). The characteristics also vary from mill to mill and thus no unambiguous conclusions regarding quality all the pulp mill effluents can be drawn.

**Paper making**

The specific freshwater consumption of a paper mill (and, by implication, the effluent volume generated) is strongly dependent on the paper grade produced and on the technical age of the paper machine (Table 3.11), as well as the availability and price of freshwater. In general, mill water is needed for showers in the paper machine wire and press section, dilution of chemicals, and process water makeup for level control in the tanks (Table 3.12).

**Paper mills using chemical or mechanical pulp (virgin fibre)**

As mentioned earlier the water consumption and effluent quality depend on, amongst other things, the age of the mill. The non-integrated wood-free fine
Table 3.9 Effluents from a modern softwood kraft pulp mill (Myrèn, 1993)

<table>
<thead>
<tr>
<th>Source</th>
<th>Flow (m³/ADt)</th>
<th>COD (kg/ADt)</th>
<th>AOX (kg/ADt)</th>
<th>P (kg/ADt)</th>
<th>Colour (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wood room (debarking)</td>
<td>2</td>
<td>5</td>
<td>9</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>Fibre line</td>
<td>1</td>
<td>5</td>
<td>20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bleach plant</td>
<td>30</td>
<td>25</td>
<td>1.0</td>
<td>36</td>
<td>40</td>
</tr>
<tr>
<td>Recovery system</td>
<td>2</td>
<td>5</td>
<td>8</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>Spills etc.</td>
<td>5</td>
<td>8</td>
<td>45</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>From mill</td>
<td>40</td>
<td>48</td>
<td>1.0</td>
<td>45</td>
<td>100</td>
</tr>
<tr>
<td>After AS treatment</td>
<td>40</td>
<td>22</td>
<td>0.5</td>
<td>20</td>
<td>100</td>
</tr>
</tbody>
</table>

ADt, air dried tonne.

paper mill, with a production of over 1 million tonnes p.a., shown in Table 3.13, represents an example of modern paper making technology with low effluent pollutant levels and low water consumption. The range of emissions can be extremely broad, as shown in the case of the tissue mill also shown in Table 3.13. Both of these paper grades use bleached chemical pulp as their major component.

Chemical pulp is much cleaner than mechanical pulp as a raw material due to the many washing stages the pulp undergoes before entering the paper machine water system. Mechanical pulp transports, amongst other things, significant amounts of pollutants, mainly so-called “anionic trash”, to the paper machine water circuit. The anionic trash is detrimental, causing operational problems in the paper machine and reducing the efficiency of additives such as retention aids (reagents added to aid the retention of constituents in the paper). In the case of peroxide-bleached Norway spruce (Picea abies) TMP, most of the anionic trash comprises anionic galacturonic acid-rich hemicelluloses, i.e. polygalacturonic acids (Thornton et al., 1993).

In mechanical pulp production the effluents result mainly from various thickening processes (e.g. thickener and pressing filtrates) indicated as water sources in Fig. 3.19. These water fractions are usually collected to form pulp mill white water (circulation water). This water typically consists of wood-originating substances (such as resin and fatty acids and other lipophilic extractives), lignin, sugars, polysaccharides, simple organic acids and salts. The concentrations are usually lower in the white water than in the different filtrates. Some matrix characteristics are shown in Table 3.14, which indicates the effect of peroxide bleaching, most often used for mechanical pulp, on the levels of dissolved and colloidal substances (DCS).

In a modern integrated (mechanical) pulp and paper mill (Fig. 3.19) the freshwater is taken in as wire section, this being the flat belt of metal or plastic mesh on which the paper web is dewatered (point B in Fig. 3.19), shower water. The water removed from the web in the paper machine is collected in the wire pit (point C in Fig. 3.19), and forms the white water from the paper machine. In most of the modern paper mills the white water is treated with a disc filter. In older mills, and especially in Central Europe, white water is treated by microflotation. The aim of the treatment is to recover fibre and to remove
Table 3.10  Different pulping effluents and their flows and quality (EIPPCB, 2001)

<table>
<thead>
<tr>
<th>Process</th>
<th>Effluent volume/flow (m$^3$/ADt)</th>
<th>BOD$_5$ (kg/ADt)</th>
<th>COD (kg/ADt)</th>
<th>AOX (kg/ADt)</th>
<th>TSS (kg/ADt)</th>
<th>Tot-N (kg/ADt)</th>
<th>Tot-P (g/ADt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Debarking</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wet debarking and press</td>
<td>3–10</td>
<td>5–15</td>
<td>20–30</td>
<td>n/a</td>
<td>n/a</td>
<td>25–35</td>
<td></td>
</tr>
<tr>
<td>Dry debarking and press</td>
<td>0.5–2.5</td>
<td>0.5–2.5</td>
<td>1–10</td>
<td>n/a</td>
<td>n/a</td>
<td>10–20</td>
<td></td>
</tr>
<tr>
<td>Kraft mill</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbleached pulp</td>
<td>20–80</td>
<td>1–20</td>
<td>7–50</td>
<td>0.2–15</td>
<td>0.1–1</td>
<td>3–40</td>
<td></td>
</tr>
<tr>
<td>Bleached pulp</td>
<td>30–110</td>
<td>0.2–40</td>
<td>4–90</td>
<td>0.2–10</td>
<td>0.1–0.8</td>
<td>5–90</td>
<td></td>
</tr>
<tr>
<td>Sulphite pulping</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bleached and unbleached</td>
<td>40–100</td>
<td>0.5–75</td>
<td>10–190</td>
<td>1.7–10</td>
<td>0.18–1</td>
<td>15–150</td>
<td></td>
</tr>
<tr>
<td>Mechanical pulping</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Groundwood</td>
<td>5–15</td>
<td>8.5–10</td>
<td>20–30</td>
<td>n/a</td>
<td>0.08–0.1</td>
<td>20–25</td>
<td></td>
</tr>
<tr>
<td>TMP</td>
<td>4–10</td>
<td>13–22</td>
<td>50–80</td>
<td>n/a</td>
<td>0.1–0.13</td>
<td>30–40</td>
<td></td>
</tr>
<tr>
<td>CTMP</td>
<td>15–50</td>
<td>17–30</td>
<td>60–100</td>
<td>n/a</td>
<td>0.11–0.14</td>
<td>35–45</td>
<td></td>
</tr>
<tr>
<td>Bleached CTMP</td>
<td></td>
<td>25–0</td>
<td>80–130</td>
<td>n/a</td>
<td>0.13–0.4</td>
<td>50–60</td>
<td></td>
</tr>
</tbody>
</table>

n/a, not available; ADt, air dried tonne.
Table 3.11  Specific freshwater consumption in modern paper mills (Sundholm, 2000; Weise et al., 2000)

<table>
<thead>
<tr>
<th>Paper grade in general</th>
<th>Typical example</th>
<th>Water consumption (m$^3$/t)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Newsprint</td>
<td>Pulp mill and newsprint line</td>
<td>5–15</td>
</tr>
<tr>
<td>Wood-free fine paper</td>
<td></td>
<td>9</td>
</tr>
<tr>
<td>Supercalandered (SC) paper</td>
<td></td>
<td>5–10</td>
</tr>
<tr>
<td>Lightweight coated (LWC) paper</td>
<td></td>
<td>10–15</td>
</tr>
<tr>
<td>Tissue</td>
<td>Paper machine and process</td>
<td>11</td>
</tr>
<tr>
<td>Liner and fluting</td>
<td>Pulp and testliner mill</td>
<td>5–15</td>
</tr>
<tr>
<td>Multiple board</td>
<td>Board line</td>
<td>10</td>
</tr>
</tbody>
</table>

Table 3.12  Freshwater consumption in papermaking (Edelmann, 1999a; Haavanlammi, 1999)

<table>
<thead>
<tr>
<th>Unit operation water use, m$^3$/t</th>
<th>Printing paper grades</th>
<th>Newspaper grades</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total fresh water consumption</td>
<td>16–26</td>
<td>27</td>
</tr>
<tr>
<td>Paper machine showers</td>
<td>6–7</td>
<td>5</td>
</tr>
<tr>
<td>Dilution/preparation of chemicals</td>
<td>2–3</td>
<td>3</td>
</tr>
<tr>
<td>Makeup waters in pulp production</td>
<td>2–3</td>
<td></td>
</tr>
<tr>
<td>Cooling of process equipment</td>
<td>3–10</td>
<td>10</td>
</tr>
<tr>
<td>Sealing waters</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>Power plant</td>
<td>2–3</td>
<td></td>
</tr>
<tr>
<td>Miscellaneous (washing, sealing etc.)</td>
<td></td>
<td>3</td>
</tr>
</tbody>
</table>

suspended solids from the water. The modern disc filters produce three filtrates; cloudy, clear and superclear filtrate. The cloudy filtrate is usually directed back to the stock preparation, but the clear and superclear filtrates are either reused – e.g. as wire section shower water – or discarded as effluents, depending on their suspended solids content and on the paper grade produced. In the worst case the clear filtrate can constitute more than a half of the total effluent load of a paper mill. The main prerequisite for recycling the clear or the superclear filtrate in the process is that they are practically free of suspended solids.

The clear filtrate from the paper machine consists of suspended solids, dissolved and colloidal substances originating from wood, and salts. In addition, the clear filtrate also contains traces of all the paper making chemicals added to the process. If the paper machine produces coated paper grades, the traces of the coating colour ingredients are also found in the clear filtrate. These include various pigments and latices. It should be noted that white water quality varies significantly between machines and, in some cases, no significant difference is seen between the clear and superclear filtrates.
Table 3.13  Consumption and emission levels of the biggest non-integrated wood-free fine paper mill in Europe and a typical tissue mill (EIPPCB, 2001)

<table>
<thead>
<tr>
<th></th>
<th>Fresh water demand (m³/t)</th>
<th>Wastewater flow (m³/t)</th>
<th>COD (kg/t. mg/l)</th>
<th>BOD₅ (kg/t. mg/l)</th>
<th>AOX (g/t. mg/l)</th>
<th>TSS (kg/t. mg/l)</th>
<th>N (g/t. mg/l)</th>
<th>Tot-P (g/t. mg/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wood-free line paper</td>
<td>6</td>
<td>4.5</td>
<td>0.44</td>
<td>0.11</td>
<td>7.</td>
<td>0.14</td>
<td>41.9</td>
<td>3.0</td>
</tr>
<tr>
<td>Tissue mill</td>
<td>7–100</td>
<td>6–100</td>
<td>2–6ᵃ</td>
<td>1–2ᵃ</td>
<td>5–15ᵃ</td>
<td>1–3ᵃ</td>
<td>6–100ᵃ</td>
<td>1–30ᵃ</td>
</tr>
</tbody>
</table>

ᵃ After wastewater treatment plant.

Figure 3.19  Water circulation system for a modern integrated mechanical pulp and paper mill showing water sources and sinks. The numbers indicate the pulp concentration in different points of the process. A = shower water taken into the wire section of the paper machine, B = wire section, C = wire pit (courtesy of K. Edelmann (Edelmann, 1999b))

Paper mills using recovered fibre
Recovered paper (RCF) has become more popular as a raw material in paper making, especially in the regions where the population density is high and paper collection systems are working efficiently. For example, out of the 6.5 Mt of paper and board manufactured in the UK in 1998, 4.7 Mt was raw material
Table 3.14 Characteristics of mechanical pulp (thermomechanical pulp, TMP, (Thornton, 1993)) and mechanical pulp mill water fractions: groundwood mill circulation water (Huhhilo et al., 2002) and plug screw feeder pressate from semi-chemical mechanical pulp mill (SCMP) (Dal-Cin et al., 1995)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Groundwood mill circulation water&lt;sup&gt;a&lt;/sup&gt;</th>
<th>SCMP mill pressate&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Thermomechanical pulp&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Average</td>
<td>Range</td>
<td>Plug screw feeder pressate</td>
</tr>
<tr>
<td>Temperature</td>
<td>°C</td>
<td>63</td>
<td>50-70</td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td></td>
<td>5.3</td>
<td>4.8-6.1</td>
<td>5.7</td>
</tr>
<tr>
<td>Conductivity</td>
<td>µS/cm</td>
<td>905</td>
<td>4.2-6.1</td>
<td>5100</td>
</tr>
<tr>
<td>Cationic demand</td>
<td>µeq/l</td>
<td>364</td>
<td>222-555</td>
<td>1200</td>
</tr>
<tr>
<td>Colour</td>
<td>PTCO</td>
<td>460</td>
<td>250-1300</td>
<td>1700</td>
</tr>
<tr>
<td>Suspended solids</td>
<td>mg/l</td>
<td>1400</td>
<td>1100-1750</td>
<td>5670</td>
</tr>
<tr>
<td>Dissolved solids</td>
<td>mg/l</td>
<td>500</td>
<td>300-750</td>
<td>2400</td>
</tr>
<tr>
<td>Soluble COD</td>
<td>mg/l</td>
<td>220</td>
<td>150-320</td>
<td>420</td>
</tr>
<tr>
<td>Total organic carbon</td>
<td>mg/l</td>
<td>620</td>
<td>190-540</td>
<td>2680</td>
</tr>
<tr>
<td>Dissolved organic carbon</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Extracted UV250 nm</td>
<td>abs.</td>
<td>2.6</td>
<td>1.7-3.7</td>
<td>37</td>
</tr>
<tr>
<td>Lignin-like COD</td>
<td>mg/l</td>
<td>300</td>
<td>190-540</td>
<td>67</td>
</tr>
<tr>
<td>Lignin + other aromatics COD</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Lignin soluble</td>
<td>mg/l</td>
<td>620</td>
<td>2680</td>
<td>209</td>
</tr>
<tr>
<td>Lignin insoluble</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Sugar COD</td>
<td>mg/l</td>
<td>300</td>
<td>190-540</td>
<td>2680</td>
</tr>
<tr>
<td>Carbohydrates</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Lipophilic extractives</td>
<td>mg/l</td>
<td>300</td>
<td>190-540</td>
<td>2680</td>
</tr>
<tr>
<td>Resin and fatty acids</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Neutral hydrophobics</td>
<td>mg/l</td>
<td>300</td>
<td>190-540</td>
<td>2680</td>
</tr>
<tr>
<td>Acidic hydrophobics</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Lignans</td>
<td>mg/l</td>
<td>300</td>
<td>190-540</td>
<td>2680</td>
</tr>
<tr>
<td>Volatile acids</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>mg/l</td>
<td>300</td>
<td>190-540</td>
<td>2680</td>
</tr>
<tr>
<td>Formic acid</td>
<td>mg/l</td>
<td>16</td>
<td>11-26</td>
<td>50</td>
</tr>
<tr>
<td>Ca&lt;sup&gt;2+&lt;/sup&gt;</td>
<td>mg/l</td>
<td>19</td>
<td>11-28</td>
<td>10</td>
</tr>
<tr>
<td>Cl&lt;sup&gt;-&lt;/sup&gt;</td>
<td>mg/l</td>
<td>6</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>SO&lt;sub&gt;4&lt;/sub&gt;&lt;sup&gt;2+&lt;/sup&gt;</td>
<td>mg/l</td>
<td>246</td>
<td>153-395</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Analysed from mill samples.
<sup>b</sup> Analysed from supernatant resulting from pulp agitated in clean water (60°C) and centrifuged in laboratory.
<sup>c</sup> DCS, dissolved and colloidal substances.

supplied by waste paper merchants (British Recovered Paper Association, 2002). For comparison, out of the 14.4 million tonnes of paper and board manufactured in Finland in 2001, 0.7 Mt of recovered fibre used as raw material equates to 5%. However, 90% of the paper produced in Finland is exported and, within Finland, around two-thirds of the paper and board used is recovered, i.e. 14 kg per capita per annum. The global average for paper and board recovery and reuse is around 40% (Finnish Forest Industries Federation, 2002).
Recovered paper loads the water circuit of paper mills with a variety of adhesives. Colloidal, dissolved, finely dispersed or water-soluble adhesives originating from recovered fibre are called secondary stickies because they agglomerate only later in the finished stock through chemical reaction, or else they appear only on the paper machine thus causing severe problems. In addition, the alkaline de-inking procedure and a high temperature promote dissolution of adhesives in the circulation water (Zippel, 2001). It has been estimated that the total amount of adhesives in DIP (de-inked pulp) is 7 kg/t of paper, 70 times the maximum permitted level of sticky material allowed in white paper manufactured from DIP (0.1 kg/t; Zippel, 2001). Volume and quality data for water from stock preparation of different paper grades using recovered fibre are shown in Table 3.15.

### 3.2.4 Current water and effluent purification systems and governing legislation

Driving forces for effluent treatment in the pulp and paper industry can be categorised as follows:

- Environmental legislation
- Better economy
- Local demands, for example lack of water resources
- Customer demand

The environmental legislation in the pulp and paper industry is likely to undergo profound changes within the next few years. The EU Directive on Integrated Pollution Prevention and Control (IPPC) includes a BAT reference document, BREF, which refers to the pulp and paper industry. This came into force in 1999 for greenfield mills, and will apply to existing mills from 2007.

| Table 3.15 Emission levels for stock preparation of different paper grades using recovered paper (EIPPCB, 2001) |
|---|---|---|---|
| Recovered paper quality | Water flow (m³/t paper) | TSS (mg/l) | COD (kg/t, mg/l) | AOX (g/t, mg/l) |
| Packaging paper | Sorted mixed paper and boars, recovered paper from stores | 0–4 | Below 200 | 27–36, 6750–9000 | < 4, 1 |
| Newsprint | De-inkable recovered paper (50:50 newsprint and magazines) | 8–16 | Below 200 | 17–27, 1700–2700 | < 10, 1 |
| LWC/SC paper | De-inkable recovered paper (50:50 newsprint and magazines) | 8–16 | Below 200 | 17–27, 1700–2700 | < 10, 1 |
| Tissue paper and market pulp | De-inkable recovered paper (50:50 newsprint and magazines); wood-free office recovered paper | 8–16 | Below 200 | 26–35, 2600–3500 | < 10, 1 |
BREF stipulates, amongst other things, permitted effluent loads and, for the first time, effluent flows for different kinds of pulp and paper mills. The directive also includes a list of candidate techniques and practises (under BAT, best available technology, Tables 3.16 and 3.17) recognised as being able to achieve the mandatory effluent loads.

The effluent loads allowed according to the IPPC directive are extremely rigorous. In particular, target levels for nitrogen and phosphorus cannot be achieved by use of conventional external (end-of-pipe) biological treatment alone. The allowed effluent flows, on the other hand, necessitate counter current water circulation systems and internal water recycling, also stated in the IPPC directive and aimed at minimising freshwater consumption.

The focus of the internal water purification used today has been mainly on the removal of suspended solids by mechanical filtration (e.g. disc filter), flotation or chemical precipitation. Flotation (or flotation combined with sand filtration, so-called flotation filtration) has been successfully used to remove suspended solids and to recover fibres, fillers and fines from the white water. However, flotation tanks generally demand a lot of floor space and the process is fairly energy intensive. The cost and the filtrate quality are very dependent on the chemicals used, which are usually expensive. Flotation is used to a large extent in mills using recovered paper as raw material. Chemical precipitation is also used within the paper machine white water treatment system. The aim is to improve the first-pass retention in the wire section and, at the same time, the suspended solids content is reduced in the white water. However, chemicals are usually expensive and the doses required can be large. Moreover, overdosing of

<table>
<thead>
<tr>
<th>Table 3.16</th>
<th>BAT requirements for mechanical pulp mills (EIPPCB, 2001)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mill type</td>
<td>Flow (m³/t)</td>
</tr>
<tr>
<td>Non-integrated CTMP mills (contribution of pulping only)</td>
<td>15–20</td>
</tr>
<tr>
<td>Integrated mechanical pulp and paper mills (e.g. newsprint, LWC and SC paper mills)</td>
<td>12–20</td>
</tr>
</tbody>
</table>

* LWC, lightweight coated; SC, supercalendered.

<table>
<thead>
<tr>
<th>Table 3.17</th>
<th>BAT demands for pulp and paper mills using chemical pulp (EIPPCB, 2001)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paper type</td>
<td>Flow (m³/t)</td>
</tr>
<tr>
<td>Uncoated fine paper</td>
<td>10–15</td>
</tr>
<tr>
<td>Coated fine paper</td>
<td>10–15</td>
</tr>
<tr>
<td>Tissue</td>
<td>10–15</td>
</tr>
</tbody>
</table>
chemicals can interfere with the sensitive chemical equilibrium in the wet processing.

Suspended solids-free process water or effluent is not always sufficiently clean for recycling purposes. When the freshwater consumption is reduced the water circulation system becomes enriched in dissolved and colloidal substances (DCS), which causes several problems within the process operation and/or in the paper product. High levels of dissolved organics result in enhanced bacterial growth in the system, possibly producing odour problems in the paper product and increasing BOD and colour levels in the effluent streams. Enrichment of multivalent ions also produces problems of scaling (carbonates and silicates) and corrosion (sulphates, chlorides, Fe$^{2+}$ and Al$^{3+}$) which tend to arise after extended periods of operation under conditions of low freshwater use. Also, brightness reversion can be caused by dissolved inorganic materials such as Fe$^{2+}$. Problems of enrichment of the DCS in the white water system can be substantially ameliorated through advanced purification methods, such as ultrafiltration (UF), as will be shown later.

The stress imposed on water resources has forced the mills to seek ways to efficiently treat their effluents to freshwater quality levels of purity. For example, the total mill effluents treated by biological and membrane processes (Bentley, 1999; Webb, 1999) or evaporation (Stevenson, 1992) have been successfully used to supplement freshwater supplies. However, coating colour effluents have proved problematic to treat because they are not degraded by biological processes and, as a result of this, chemical precipitation has been widely employed. This produces a solid waste that must be landfilled. During the 1990s concentration of coating colour effluents by UF became more common, mainly for economic reasons: UF costs have decreased whereas landfill disposal costs have increased. In the IPPC directive UF treatment of the coating colour effluents is now stated as a BAT technology.

### 3.2.5 Membranes in the pulp and paper industry

There has been increased interest in membrane filtration applications in the pulp and paper industry over the last 15–20 years, for reasons already stated above as well as in Section 1. Membrane processes offer a high level of purification coupled with a low footprint and relatively low energy consumption, especially when compared with the competing desalination technology of evaporation.

**Existing full-scale membrane plants**

The first full-scale reverse osmosis plant was installed in the white water system of a board machine of Green Bay Packaging Inc. in the USA in 1974 (Macleod, 1974). Later when the water balance of the board machine changed the RO plant became redundant. Since the 1980s tubular module ultrafilters have been successfully adopted for such purposes as concentration and fractionation of spent sulphite liquor (Anon., 1982; PCI Membrane Systems, 1988), deresination (Paterson Candy, 1987) and bleaching effluent treatment (Haagensen, 1982; Okamoto et al., 1985; Jönsson, 1987; Wickström, 1997). An early example of
the use of ultrafiltration is the tubular ultrafiltration plant at Borregaard sulphite pulp mill in Norway, which has been used to process spent sulphite liquor since 1981 (Table 3.18). The UF concentrate contains high molar mass fractions of lignosulphonates and small amounts of sugar and salts. The concentrate is used for vanillin production.

To meet the emission levels stated by legislation or to improve the efficiency of the external biological treatment some specific effluents, e.g. bleaching effluents with high COD loads, have been separately pre-treated prior to biotreatment. For these reasons several ultrafiltration plants were installed in the early 1980s to treat the first alkaline stage effluent, E1, from the bleaching process (Haagensen, 1982; Okamoto et al., 1985; Jönsson, 1987). For example, in the Taio Paper Co. mill the COD reduction obtained was 79%, equating to 5.5 t d⁻¹. This was enough to meet the emission level and the efficiency of the biological effluent treatment plant was significantly increased. The concentrate was incinerated with the black liquor. The average permeate flux of the polysulphone UF membranes used for the 3.5 million litres per day (MLD) plant was about 100 \( \text{L m}^{-2} \text{h}^{-1} \) (LMH), giving a total membrane area requirement of 1480 m².

Even though the processes using tubular membranes perform well, they do so at a substantial cost. The packing density of the tubular membranes is relatively low (Table 2.5) such that membrane plants based on this modular configuration have a relatively large footprint. In addition, fluxes attainable from tubular membranes are limited by the high fouling propensity of the liquids arising.

The cross-rotational (CR) filter (Section 2.1.4, Fig. 2.12), originally developed by ABB Flootek, Sweden (now Metso PaperChem, Finland), entered the market in the late 1980s. It was shortly adapted for mill-scale ultrafiltration applications, such as treatment of bleaching effluents and board machine white water as well

### Table 3.18 Ultrafiltration of spent sulphite liquor at Borregaard Industries, Norway (PCI Membrane Systems, 1988)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Details/value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Membrane</td>
<td>BX1, PCI Membrane Systems</td>
</tr>
<tr>
<td>Total membrane area, m²</td>
<td>1120</td>
</tr>
<tr>
<td>Process design</td>
<td>2 lines with 6 stages in series with diafiltration</td>
</tr>
<tr>
<td>Feed flow, m³ h⁻¹</td>
<td>50</td>
</tr>
<tr>
<td>Concentrate flow, m³ h⁻¹</td>
<td>16</td>
</tr>
<tr>
<td>Solids content, %</td>
<td>Feed 12</td>
</tr>
<tr>
<td></td>
<td>Concentrate 22</td>
</tr>
<tr>
<td>Pressure, bar</td>
<td>10–15</td>
</tr>
<tr>
<td>Cross-flow velocity</td>
<td>High</td>
</tr>
<tr>
<td>Temperature, °C</td>
<td>60–65</td>
</tr>
<tr>
<td>pH</td>
<td>4.2–4.5</td>
</tr>
<tr>
<td>Washing</td>
<td>Alkaline detergent daily for 2 hours</td>
</tr>
<tr>
<td>Relative costs</td>
<td>Membranes 65%</td>
</tr>
<tr>
<td></td>
<td>Energy 26%</td>
</tr>
<tr>
<td></td>
<td>Mechanical spare parts 6%</td>
</tr>
<tr>
<td></td>
<td>Chemicals 3%</td>
</tr>
</tbody>
</table>


as concentration of coating colour. The filter normally operates at low pressures – as low as 0.8 bar – with the main energy demand being mechanical, between 3 and 5 KWh m$^{-3}$. As a stacked disc system, the module imposes a lower footprint than tubular systems whilst achieving higher fluxes.

Four CR ultrafilters were successfully used in 1988–1992 at MoDo Husum sulphate pulp mill in Sweden to treat the E1/O bleaching effluent (Jönsson and Trägårdh, 1990). The membrane area of one filter was 50 m$^2$ and its capacity 50 m$^3$ h$^{-1}$, giving a flux of 1000 LMH. The concentration factor was 15, the pressure 0.2–0.8 bar and temperature 50–60°C. Hoechst PES25 polyethersulphone UF membranes with a MWCO of 25 kDa were used. The concentrate was incinerated and the permeate was directed to the external biological treatment plant.

CR filters were also installed in 1989 at Rottneros Board Svaneholm in Sweden to treat board machine white water (Jönsson, 1990; Gavelin, 1991; Anon., 1996). Two CR1000 filters (membrane area 100 m$^2$ and membrane cut-off 100 kDa) are used to treat the white water removing 3 t d$^{-1}$ of suspended solids. The permeate is directed back to the board machine and the retentate is first concentrated in an evaporator and then burned together with the stripped bark. The content of suspended solids in the white water has been reduced from 5 to 2% and the downtime associated with machine cleaning and maintenance reduced. Also, the efficiency of the secondary treatment plant has been increased due to the lower organic loading. The energy consumption is 6–8 KWh m$^{-3}$ of permeate. Successful pilot tests with UF for white water treatment have also been carried out at Stora Kabel (Kothe and Schroth, 1999) and StoraEnso Uetersen (Kluter, 1999; Huhtamäki et al., 2002). For the UF plant the payback time calculated on the basis of the pilot trials in Stora Kabel is 4.9 years and the internal rate of return 21%. The cost for fresh water and wastewater is about $0.5 per m$^3$. Through recycling, and thus the reduction in freshwater consumption, about $0.15 per m$^3$ is saved. A recent example of the mill scale application of a CR filter to the treatment of paper mill circulation water (Teppler et al., 1998) is detailed in Section 5.6.

As with other industrial sectors, the cost benefit of the water recycling plant is substantially improved if other resources are simultaneously recovered for reuse. In paper mills producing, for example, coated high-quality printing papers, such as lightweight coated LWC paper grades, a significant amount of coating colour is discharged in the effluent. Moreover, the waste coating colour adheres to equipment, which then requires frequent cleaning. This represents a significant economical loss, as well as the colour in the effluent being an environmental issue. In many mills coating colour effluents are still treated with precipitation and the solid waste then landfilled. By employing ultrafiltration colour components in the coating colour kitchen effluent stream can be recycled, along with the water. The dilute effluents are concentrated with ultrafiltration membranes to an appropriate total solids content after which the retentate is used in the preparation of fresh coating colour. The permeate can be used for dilution or washing purposes in the coating colour kitchen. The payback time of such membrane plants is less than a year depending on the price of the coating
colour raw materials. Today, there are about 25 CR plants (Anon., 1996; Kenny, 1997; Nygåård et al., 1998; Alho et al., 1998; Sutela, 2002) and several tubular membrane plants (Camatta, 1992; Stuffer, 1996; GAW, 2002) worldwide treating coating colour effluents (Fig. 3.20).

In a zero effluent discharge (ZLD) mill producing linerboard at the McKinley Paper plant in New Mexico, a waste stream of 3.2 m$^3$ t$^{-1}$ paper produced is cleaned using membranes following extensive pretreatment including dissolved air flotation and activated sludge treatment (Bentley, 1999; Webb, 1999; Fig. 3.21). Following the activated sludge plant the clarified water is treated by continuous microfiltration (CMF) using hollow fibre (HF) modules (Vivendi Memcor) operating out-to-in. The particles or foulants, which adhere to the outside surface of the hollow fibres, are then removed by combining the backflush cycle with air sparging. It is only rigorous pretreatment that allows the use of HF modules, which would normally not be appropriate for pulp and paper applications due to the high solids loading. The MF permeate is further treated with reverse osmosis (RO) using brackish water elements, generating 0.7 m$^3$ water per tonne of paper to the mill. The membrane processes are followed by mechanical vapour recompression, MVR, evaporation, which regenerates another 0.4 m$^3$ water t$^{-1}$ of paper to the mill. This leaves only 0.8 m$^3$ water t$^{-1}$ paper to be taken in as make-up water – 25% of the total demand.

Reverse osmosis has also been used to treat wood plant effluent in the production of medium density fibreboard, or MDF (Finnemore and Hackney, 2000). The wood plant effluent is firstly flocculated and filtered in a filter press. The water is then treated with a multimedia filter prior to RO. The concentrate can be reused in production and the permeate is treated with a carbon filter before recirculation to the process main stream. The COD content of the influent to the filter press is 24 000 mg l$^{-1}$, and is reduced to 7000 mg l$^{-1}$ after the filter

Figure 3.20  Ultrafiltration plant treating coating colour effluent (Mäkinen, 2000)
press and 100 mg l\(^{-1}\) in the RO permeate. Thus the RO removes 98–99% of the COD in the original wood room effluent. In this case the RO process was found to be significantly cheaper than any other treatment methods. This was because the plant is mechanical, including only a few process stages thus reducing the capital investment cost compared to other systems such as evaporators or biological systems. Operating costs are substantially lower than those of the traditional treatment technologies as outputs can be recycled or reused thereby reducing the volumes and generating financial benefits from reduced effluent disposal and freshwater demand. In addition, there is no generation of by-products, such as waste activated sludge. Thus the costs for the RO membrane process are only around $1.5 per m\(^3\) ultimate disposal: the biological and evaporative processes would have respectively cost, in this case, £5 and £15 per m\(^3\), based on ultimate disposal.

One of the first nanofiltration (NF) plants to be employed in the pulp and paper industry was reported by Lien and Simonis (1995). This 10 ML/D plant located in the Pacific rim for cleaning paper mill and total effluent, produced from newsprint, printing paper and de-inked pulping operations, employs spiral wound NF modules (Osmonics). As with any dense membrane process, extensive pretreatment is required (Fig. 3.22) to reduce the fouling propensity of the water. The cleaning cycle time is over two weeks.

In the 1990s another high-shear membrane module, the VSEP filter (vibrating shear enhanced processing filter; Section 2.1.4, Fig. 2.12) entered the market. Existing mill-scale applications of the VSEP filter in the pulp and paper industry include weak black liquor, box plant effluent, and MDF wastewater treatment.
Figure 3.22 Schematic flow sheet of a paper mill spiral-wound nanofiltration process (Lien and Simonis, 1995), Osmonics, Inc.

(Shing et al., 1997; Nuortila-Jokinen and Nyström, 2000a, 2000b) and as well as recycle mill effluent treatment (Monroe, 1997; New Logic Int., 2002).

**Bench- and pilot-scale research**

As already mentioned, implementation of membrane technology within the pulp and paper industrial sector accelerated in the early 1990s, with the development of less expensive membrane materials, more robust modules and more sophisticated technologies such as CR and VSEP filters. A concomitant acceleration in research activity took place at this time, focused on the treatment of bleaching effluents and the closing of a bleaching plant water cycle (Jönsson et al., 1988; Jönsson and Petersson, 1989; Afonso and de Pinho, 1991; Luque et al., 1994; de Pinho et al., 1995, 1998; Rosa and de Pinho, 1995; Geraldes and de Pinho, 1995; Fält et al., 1999, 2000, 2001a, 2001b). Another area of interest has been the filtrates and circulation waters originating from mechanical pulp production (Beaudoin et al., 1992; Paleologou, 1994; Salovius et al., 1993; Konishi et al., 1998; Edelmam, 2001), and paper mill white water and total effluent have also been intensively studied (Nuortila-Jokinen et al., 1995a, 1995b; Nuortila-Jokinen and Nyström, 1996; Mänttäri and Takeshita, 1997; Nuortila-Jokinen, 1997; Mänttäri et al., 1997, 1998; Kuide and Yamamoto, 1999; Mänttäri et al., 1999; Huuhilo et al., 2001). Studies have also been conducted on the concentration of coating colour effluents (Jönsson et al., 1996; Singh et al., 1997, 1999; Takeshita, 1997; Kuide and Yamamoto, 1999; Nuortila-Jokinen and Nyström, 2000).

Pulp and paper mill process waters and effluents have a very high fouling propensity, such that the most effective membrane material for these applications seem to be highly hydrophilic, almost uncharged regenerated cellulose. For the coating colour effluents, on the other hand, the more hydrophobic membranes are preferred (Singh et al., 1997, 1999; Nuortila-Jokinen and Nyström, 2000).
In addition to the membrane material, the module structure and the operating parameters have a significant impact on membrane process performance. In general, the higher the shear rate at the membrane surface, the higher the flux and the lower the fouling rate. The cross-flow velocity should exceed 4 m s\(^{-1}\) (Jönsson and Wimmerstedt, 1985) in a tubular membrane and 3.2 m s\(^{-1}\) in flat sheet module to avoid fouling (Mänttäri et al., 1998). As already stated (Section 2.1.4), in the high-shear modules such as the CR and the VSEP filters, the rotation speed and the vibration amplitude have a direct impact on the flux (Figs. 3.23 and 3.24).

The concept of critical flux (Section 2.2.4) has been introduced for nanofiltration of pulp and paper waters by Mänttäri and Nyström (2000), with respect to the spiral-wound DS-5DK membrane (Osmonics). The efficacy of operation below the critical flux has been clearly demonstrated in the M-Real Kirkniemi case described in Section 5.6. Because the CR filters are operated at 0.8 bar, which yields fluxes well below the critical flux (depends on the membrane and we have not measured it in the CR only in the VSEP and C30F) for the membranes, hardly any fouling has arisen for this plant. The membranes are washed only once a week as a part of the routine operational procedure, rather than on demand. The VSEP filter operates with almost no fouling in nanofiltration applications (Fig. 3.25) whereas ultrafiltration membranes are usually readily fouled in the VSEP filter.

The main foulants arising from diagnosis of membranes that have been used for the treatment of mechanical pulp and paper mill waters are resins, fatty acids and other wood extractives, sugars, and metal ion coagulated organic matter (Carlsson et al., 1998; Ramamurthy et al., 1995; Nuortila-Jokinen, 1997; Huuhilo et al., 2001). The adsorption of such materials is increased at higher membrane hydrophobicities (Fig. 3.26).

![Figure 3.23](image)

**Figure 3.23** Effect of rotor blade tip speed in the CR filter on the permeate flux for the filtration of acidic clear filtrate from a paper machine. NF200 membrane from Dow-Film Tec, pH 4.9, 35°C, 10 bar TMP
The membrane surface charge along with that of the feed water constituents has a great influence on membrane process operation. As with much natural organic matter, most of the substances of wood origin in paper mill effluents are negatively charged at higher pH through dissociation yielding, for example, phenolic or carboxylate organic anions. On the other hand, the process pH in paper making is either acidic (pH 4–5) or neutral (pH 7–8). At the lower pH values solutes become associated, such that electrostatic repulsion between them and the membrane surface may be substantially decreased. As seen in Fig. 3.27, at neutral pH values (nCF, neutral clear filtrate, pH 7) the flux remains more or less stable, but in acidic solutions (aCF, acid clear filtrate, pH 5) the flux declines steeply with concentration (or volume reduction) factor because of the increased acidity of the retentate with increasing retentate concentration. As the volume reduction factor approaches 8, the flux decreases to zero.

Compounds originating from the wood itself are not as onerous to the membrane purification process as the chemicals added to the manufacturing process, even though they are found at very low concentrations in the paper mill waters. This is probably due to the fact that many of these chemicals carry a predominantly positive charge or else are surface active. In addition, coating colour pigments, such as clay (kaolin), which enter the paper machine water circuit through the use of broke (i.e. paper makers own waste paper created during the paper making process) have been observed from electron microscopic analysis to form dense layers on the membrane surface. Cationic retention aids or fillers, such as cationic starch, also tend to adsorb on the membrane to form a gel-like layer under pressure.
Figure 3.25 Nanofiltration of paper machine clear filtrate with PVD-1 membrane in a VSEP L-filter. Membrane area 465 cm², pH 7, amplitude 32 mm, temperature 30–40°C. The fluxes are recalculated to 35°C. PWFₐ = pure water flux before filtration, PWFₐ = pure water flux after filtration (Huuhilo et al., 2001)

Figure 3.26 Amounts of different extractives found in the membranes C30F, PA 50H, and PES 50H after UF of grinding room circulation water (Huuhilo et al., 2001)

The particle shape has a significant impact on membrane flux (Fig. 3.28a). Microfiltration of the coating colour components with a 0.1 μm Teflon Typar (Tetratec) membrane reveals that the round calcium carbonate (CaCO₃) particles (usually ground limestone or marble) yield a considerably higher flux than plate-like particles such as clay and talc. Further, the hydrophobic talc gives a higher flux than clay, whose particles tend to be positively charged under
the normal paper process operating conditions. This is due to the hydrophobic–hydrophobic interaction between the talc and the Teflon membrane; i.e. repulsion between the hydrophobic particle and surface. However, when these particles are mixed with other coating colour raw materials, the effect of charge and hydrophobicity are greatly ameliorated, presumably because of aggregation and adsorptive reactions between of particles and solutes (Fig. 3.28b).

As with almost all industrial effluents, the variability of the water quality presents a significant challenge to the membrane material and module. This demands that the membrane technology must be robust to extreme conditions of, in particular, temperature and pH, as well as being tolerant of shock loads. It
can also be beneficial to include pretreatment to remove key foulants. Various biological treatment processes have been assessed for this duty, and have proved to be very effective in removing low molar mass organic substances from the feed water (Pauly, 2000; Tardif and Hall, 1997; Dufresne et al., 1998; Nuortila-Jokinen et al., 1999; Suvilampi et al., 1999; Huuhilo et al., 2002). Foulants can also be chemically bonded to form flocs through the addition of flocculating chemicals, such as retention aids, fixatives and coagulants, thus preventing them from interacting with the membrane (Nuortila-Jokinen et al., 1998). In addition, ozonation (Nuortila-Jokinen and Nyström, 2001) and electrodialysis (Geraldes and de Pinho, 1995) have also been trialled.

3.2.6 Conclusions

The pulp and paper industry is a complex one with many different kinds of mills, products and processes. Effluents therefore vary significantly in quality depending on the process from which they originate. Due to this complexity the requirements of water and effluent treatment at the different points of the manufacturing process vary significantly. However, it is evident that there is a great demand for an effective separation process, not the least due to the increasing rigour of environmental legislation and its enforcement.

Membrane technologies have been demonstrated as being appropriate for many water recycling duties within the pulp and paper industry. Existing treatment schemes for concentration and fractionation of spent sulphite liquor, deresination and bleaching effluent treatment by ultrafiltration include both conventional tubular modules and high-shear systems. Membranes are particularly advantageous when recovery of both chemical reagents and clarified water is possible. It is likely that future membrane materials and modules tolerant of temperatures close to 100°C and providing high fluxes with low fouling propensity will be developed, and that the ongoing downward pressure on membrane costs will continue to increase the economic competitiveness of this technology for water recycling applications within the industry.

Acknowledgements

The authors wish to thank the Academy of Finland for financial support. Colleagues at the Laboratory of Membrane Technology and Technical Polymer Chemistry at Lappeenranta University of Technology are warmly thanked for their co-operation over the years. Membrane and module as well as chemical manufacturers are acknowledged for kindly giving us their products to test.

References


3.3 The textile industry

3.3.1 Categories of textile processing operations

Textile industry processes comprise those which convert natural (e.g. cotton, wool, silk, etc.) and synthetic (e.g. viscose, polyester, acrylic) fibres into fabrics and other products. Four key activities can be identified within this industrial sector (Mattioli et al., 2002):

- the treatment of raw materials (preparation/production of textile fibres/yarns),
- the production of knitted/woven fabrics,
- the finishing of fabrics (i.e. changing some physical property of the fabric to meet the end use requirement), and
- production of products (e.g. garments, carpets, etc) from the fabric.

In 1998 world trade in textiles was worth approximately $370 billion, or about 6.3% of global merchandise trade (WTO, 1998). USA textile exports account for $19 billion and imports, principally from Mexico and China, around $77 billion. Around 60% of textile production takes place in Europe (29%) and the Americas, with most of the remaining production taking place in Asia (Stengg, 2001). Within the European Union, which is characterised by a large number of small-to-medium enterprises, Italy accounts for 31% of all textile and clothing manufacturing activities – more than double that of the UK (15%), Germany (14%) or France (13%). Most of this activity is accounted for by clothing manufacture.

A number of textile manufacturing processes are chemical wet processing operations necessary to properly prepare, purify, colour or finish the product. This results in the production of wastewater whose pollution load arises not only from the removal of impurities from the raw materials but also from the residual chemical reagents used for processing. The freshwater demand is specific to the type of textile processing operation, the type of material or final product and the specific machine or technique used. However, the water demand for wet processing operations is invariably high (Table 3.19), more than 5000 m³ day⁻¹ for a large mill. The industry is thus perceived as generating large volumes of effluent which are extremely variable in composition and pollution load, the variability arising from the diversity in the types of transformation processes used and the wide range of chemicals involved.

Identifying suitable pollution abatement or water recycling technologies is made difficult by the combining of effluent streams from individual operations, resulting in large variations in effluent chemical composition. Clearly, candidate waste treatment techniques need to be dedicated to individual process effluents, rather than the combined discharge, in order to be reliable and effective. However, this is made extremely difficult in real plants by the sheer number of individual processes contributing to the pollutant load on the combined effluent
Table 3.19  Typical water usage, 1 kg⁻¹ of product, in textile wet processing of woven fabrics (EPA, 1997)

<table>
<thead>
<tr>
<th>Subcategory</th>
<th>Minimum</th>
<th>Median</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simple processing</td>
<td>12.5</td>
<td>78.4</td>
<td>275.2</td>
</tr>
<tr>
<td>Complex processing</td>
<td>10.8</td>
<td>86.7</td>
<td>276.9</td>
</tr>
<tr>
<td>Complex processing plus desizing</td>
<td>5.0</td>
<td>113.4</td>
<td>507.9</td>
</tr>
</tbody>
</table>

stream. Effluent reclamation and reuse thus only becomes viable for individual wastewater streams, where the compositional variability is reduced, and/or in cases where either the discharge consents are stringent (or else the discharge costs high) or the treated effluent has some added value. Both these criteria are pertinent to dyeing wastewater streams, where the possibility exists both to recover chemicals and recycle the treated wastewater (Diaper et al., 1996).

As a rudimentary simplification the USEPA grouped the industry into nine categories in promulgating its guidelines (EPA, 1982). Table 3.20 gives effluent characteristics for the seven most important of categories, these being:

- raw wool scouring,
- yarn and fabric manufacturing,
- wool finishing,
- woven fabric finishing,
- knitted fabric finishing,
- carpet finishing, and
- stock and yarn dyeing and finishing.

It should be stressed that the figures quoted in Table 3.20 are average figures for complete processes which may entail a number of individual unit operations. Since many textile processing operations are batch, there are broad temporal variations in effluent quality. Variations also arise even within specific individual operations due to the different designs of the actual technology being used. The selection of suitable strategies for pollution abatement and/or water recycling even for specific unit operations is therefore not straightforward, and has to be considered on a case-by-case basis. On the other hand, and in common with most industrial effluent recycling problems:

- recycling is simplified by segregation of the various waste streams, and
- membrane technologies offer the most promise of all candidate treatment processes on the basis of the treated water quality being largely independent of the feedwater quality.

3.3.2 Effluents from textile processing unit operations

The complete textile manufacturing process involves a number of individual unit operations, each generating effluents of substantially different qualities. For
Table 3.20  Textile processing categories and effluent characteristics (EPA, 1978, 1997)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Categories*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>BOD/COD</td>
<td>0.2</td>
</tr>
<tr>
<td>BOD (mg/l)</td>
<td>6000</td>
</tr>
<tr>
<td>TSS (mg/l)</td>
<td>8000</td>
</tr>
<tr>
<td>COD (mg/l)</td>
<td>3000</td>
</tr>
<tr>
<td>Oil and grease (mg/l)</td>
<td>5500</td>
</tr>
<tr>
<td>Total chrome (mg/l)</td>
<td>0.05</td>
</tr>
<tr>
<td>Phenol (mg/l)</td>
<td>1.5</td>
</tr>
<tr>
<td>Sulphide (mg/l)</td>
<td>0.2</td>
</tr>
<tr>
<td>Colour (ADMI)b</td>
<td>2000</td>
</tr>
<tr>
<td>pH</td>
<td>8.0</td>
</tr>
<tr>
<td>Temperature (“C)</td>
<td>28</td>
</tr>
<tr>
<td>Water usage (l/kg)</td>
<td>Minimum</td>
</tr>
<tr>
<td></td>
<td>Median</td>
</tr>
<tr>
<td></td>
<td>Maximum</td>
</tr>
</tbody>
</table>

* Categories description:

b  ADMI (American Dye Manufacturers Institute) colour values result from a special procedure for determination of colour in dyeing wastewaters (Allen et al., 1972; Little, 1978).

e.g., a woven cotton fabric finishing mill (Fig. 3.29) will typically have processes for preparing the material (which will include some or all of the processes of singeing, desizing, scouring/washing, mercerising and bleaching), colouring it (dyeing and printing) and then fixing these colouring reagents mechanically and/or chemically. The process can thus generate up to six different liquid effluents which are conventionally combined to yield an effluent quality similar to that listed under Category 4 in Table 3.20, and discharged to sewer. However, since some components, and dyes in particular, are not readily removed by conventional municipal wastewater treatment, which is based largely on primary sedimentation followed by biological treatment, surcharges or supplementary consents may be in place for discharges of certain textile effluents. A specific UK example of discharge consents based on colour, and specifically UV absorption, imposed by the regulatory body (the Environment Agency) is given in Section 5.7.

Detailed technical descriptions of the most usual operations within the textile industry have been reported by many authors (Cooper, 1978; Nolan, 1972; OECD, 1981), along with effluent water quality data, which generally relates directly to the water at the end of the batch operation which is then discharged. However, many conventional wastewater quality determinants such as COD, TOC, TDS and TSS generally go unreported. Values for other parameters, taken from Cooper (1978), are reported in Table 3.21 along with some water consumption values. The values are subject to considerable variation arising
from differences in design of the specific process technology. For example, batch dyeing with reactive dyes, at around 38 l water per kg fabric, can demand almost 10 times as much water as continuous dyeing with vat dyes (ETBPP, 1997). The data in Table 3.21 thus relate to expected or most probable pollution loads resulting from each wet chemical unit operation in the textile manufacturing process, and do not incorporate the whole range of water qualities that may be encountered in practice. A more comprehensive listing of individual chemical components arising in specific effluent streams is given in Table 3.23.

Specific wet processes used in textile manufacturing are briefly described below. Non-wet processing techniques, such as singeing, printing, mechanical finishing, weaving and fabrication do not give rise to significant quantities of liquid effluent.

**Sizing**

In the transformation of raw materials to textile products sizing is usually the first process in which wet processing is involved. Substances such as starch, modified starch, polyvinyl alcohol, polyvinyl acetate, carboxymethyl cellulose and gums are applied to the warp in order to increase its tensile strength and smoothness. During this operation wastewater results from the cleaning of sizing boxes, rolls, size mixer and sizing area. Their volume is low but, depending on the recipe used, can contain high levels of BOD, COD and TSS (Cooper, 1978). In the case of 100% synthetic warps sizing, if used, is usually carried out with synthetic polymers. Yarns for use as knitted fabrics are treated with lubricants (mineral, vegetable or ester-type oils) or waxes rather than sizes.
Table 3.21  Pollution loads of textile wet operations (from Cooper, 1978)

<table>
<thead>
<tr>
<th>Fibre</th>
<th>Process</th>
<th>pH</th>
<th>BOD (mg/l)</th>
<th>S (mg/l)</th>
<th>Water use (l/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Desizing</td>
<td>–</td>
<td>1700–5200</td>
<td>16 000–32 000</td>
<td>3–9</td>
</tr>
<tr>
<td>Cotton</td>
<td>Scouring or kiering</td>
<td>10–13</td>
<td>50–2900</td>
<td>7600–17 400</td>
<td>26–43</td>
</tr>
<tr>
<td></td>
<td>Bleaching</td>
<td>8.5–9.6</td>
<td>90–1700</td>
<td>2300–14 400</td>
<td>3 124</td>
</tr>
<tr>
<td></td>
<td>Mercerising</td>
<td>5.5–9.5</td>
<td>45–65</td>
<td>600–1900</td>
<td>232–308</td>
</tr>
<tr>
<td></td>
<td>Dyeing</td>
<td>5–10</td>
<td>11–1800</td>
<td>500–14 100</td>
<td>8–300</td>
</tr>
<tr>
<td></td>
<td>Scouring</td>
<td>9–14</td>
<td>30 000–40 000</td>
<td>1129–64 448</td>
<td>46–100</td>
</tr>
<tr>
<td></td>
<td>Dyeing</td>
<td>4.8–8</td>
<td>380–2200</td>
<td>3855–83 15</td>
<td>16–22</td>
</tr>
<tr>
<td>Wool</td>
<td>Washing</td>
<td>7.3–10.3</td>
<td>4000–11 455</td>
<td>4830–19 267</td>
<td>334–835</td>
</tr>
<tr>
<td></td>
<td>Neutralisation</td>
<td>1.9–9</td>
<td>28</td>
<td>1241–48 30</td>
<td>104–131</td>
</tr>
<tr>
<td></td>
<td>Bleaching</td>
<td>6</td>
<td>390</td>
<td>908</td>
<td>3–22</td>
</tr>
<tr>
<td>Nylon</td>
<td>Scouring</td>
<td>10.4</td>
<td>1360</td>
<td>1882</td>
<td>50 67</td>
</tr>
<tr>
<td></td>
<td>Dyeing</td>
<td>8.4</td>
<td>368</td>
<td>641</td>
<td>17–33</td>
</tr>
<tr>
<td></td>
<td>Scouring</td>
<td>9.7</td>
<td>2190</td>
<td>1874</td>
<td>50–67</td>
</tr>
<tr>
<td>Acrylic</td>
<td>Dyeing</td>
<td>1.5–3.7</td>
<td>175–2000</td>
<td>833–1968</td>
<td>17–33</td>
</tr>
<tr>
<td></td>
<td>Final scour</td>
<td>7.1</td>
<td>668</td>
<td>1191</td>
<td>67–83</td>
</tr>
<tr>
<td></td>
<td>Scouring</td>
<td>–</td>
<td>500–800</td>
<td>–</td>
<td>25–42</td>
</tr>
<tr>
<td>Polyester</td>
<td>Dyeing</td>
<td>–</td>
<td>480–27 000</td>
<td>–</td>
<td>17–33</td>
</tr>
<tr>
<td></td>
<td>Final scour</td>
<td>–</td>
<td>650</td>
<td>–</td>
<td>17–33</td>
</tr>
<tr>
<td>Viscose</td>
<td>Scouring and Dyeing</td>
<td>8.5</td>
<td>2832</td>
<td>3334</td>
<td>17–33</td>
</tr>
<tr>
<td></td>
<td>Salt bath</td>
<td>6.8</td>
<td>58</td>
<td>4890</td>
<td>4–13</td>
</tr>
<tr>
<td>Acetate</td>
<td>Scouring and Dyeing</td>
<td>9.3</td>
<td>2000</td>
<td>1778</td>
<td>33–50</td>
</tr>
</tbody>
</table>

Desizing
Desizing removes the substance applied to the yarn in the sizing operation by hydrolysing the size into a soluble form. The methods of desizing, and therefore the wastewater characteristics, vary according to the size used (Table 3.22). Desizing can be as simple as hot washing with detergents for synthetic sizes or more complicated, for example enzyme-augmented degradation, for starch and modified starch (PRG, 1983; Nolan, 1972; OECD, 1981). The pollution load of desizing effluents results from surfactants, enzymes, acids or alkalis used in the size recipes, as well as the sizes themselves (Smith, 1989). The generated wastewater can be the largest contributor to the BOD and TS in a mill effluent (Nolan, 1972), as indicated in Table 3.21. However, if sizing is carried out using synthetic materials BOD and TSS reductions of up to 90% can be achieved on desizing (Smith, 1989).

Scouring
Scouring can be performed on both natural and synthetic materials to remove residual reagents. The intensity of the scouring process is dependent on the type of material. Cotton is scoured to remove natural waxes, pectins, spinning oils and other non-cellulosic components using hot alkaline solutions (caustic soda and soda ash) containing detergents or soaps. Herbicides, insecticides, defoliants and desiccants used in the growing of cotton, along with fungicides, such as pentachlorephenols used to prevent mildew during storage and transportation, can also arise in scouring effluents. Cotton scouring waste liquors are thus chemically aggressive and may be toxic. Their solids content,
resulting from the alkali and from impurities in the raw cotton material, is generally high (Table 3.22). Along with desizing, cotton scouring generates very high BOD concentrations. These two processes thus make by far the greatest contribution to effluent BOD in the wet phase processing of cotton goods (Nolan, 1972; Smith, 1989). When synthetic sizes are used desizing and scouring are usually carried out in a single operation. Since synthetic fibres are free from chemical impurities, 100% synthetic fabrics (woven or knitted) require only light scouring in order to remove sizes and lubricants. The process is not normally a significant source of organic or suspended solids pollution. In cases where desizing, scouring and dyeing are performed simultaneously, effluent with an increased pollution load results (Table 3.21).

Raw wool scouring is the highest-polluting operation within the textile industry (Table 3.20). The large volumes of effluent and high levels of contaminants generated by this operation have made it an area of the industry of key concern, and much work has been carried out in this area towards abatement of pollution from this process (BTTG, 1992; Nolan, 1972; OECD, 1981). The pollution load results from impurities present in the raw wool, (wax, urine, faeces, vegetable and mineral dirt, and parasite-control chemicals) together with soap, detergent and alkali used during the scouring and washing processes. The use of some of the more onerous organochlorine chemicals in sheep dipping has been restricted by legislation in recent years, but there remain chemicals such as organophosphates that are still used and so arise in raw wool scouring effluents (Shaw, 1994a,b). Due to their non-biodegradability or toxicity, many impurities in scouring effluents (Table 3.23), such as antistatic agents (synthetic fibres), pesticides, cotton waxes and wool grease or wax, can pose problems in the operation of biological treatment systems. Scouring of woollen goods is generally duplicated downstream to remove added substances. These include oils and weaving sizes or lubricants, which are removed using detergents.

**Bleaching**

Bleaching removes the natural yellow hue of cotton, increasing its whiteness. This operation is generally required if the finished fabric is to be white or dyed a light colour. It is usually carried out by chemical oxidation with sodium hypochlorite or hydrogen peroxide. Auxiliary chemicals such as sulphuric acid, hydrochloric acid, caustic soda, sodium bisulphite, surfactants and

| Table 3.22 Pollution load from desizing of 50:50 polyester/cotton (PRG, 1983) |
|-----------------------------------|--------|--------|--------|--------|
| Desizing                          | pH     | BOD (mg/l) | TSS (mg/l) | TDS (mg/l) | Oil/grease (mg/l) |
| Enzyme starch                     | 6–8    | 3078     | 6155     | 1583     | 288               |
| Polyvinyl alcohol                 | 6–8    | 200      | 400      | 4029     | 192               |
| Carboxymethyl cellulose          | 6–8    | 314      | 400      | 4349     | 751               |

Industrial waters 137
### Table 3.23 Pollutants from textile wet processing operations (Corriéa et al., 1995; data from OECD, 1981)

<table>
<thead>
<tr>
<th>Process/ fibres</th>
<th>Substances</th>
<th>Organic (biodegradability)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Inorganic</td>
<td></td>
</tr>
<tr>
<td><strong>Desizing</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cotton</td>
<td>Na⁺</td>
<td>CO₃²⁻</td>
</tr>
<tr>
<td>Linen</td>
<td>Ca²⁺</td>
<td>CO₃²⁻</td>
</tr>
<tr>
<td>Viscose</td>
<td>Na⁺</td>
<td>SO₄²⁻</td>
</tr>
<tr>
<td>Silk</td>
<td>Na⁺</td>
<td>CO₃²⁻</td>
</tr>
<tr>
<td>Acetates</td>
<td>NH₄⁺</td>
<td>PO₄³⁻</td>
</tr>
<tr>
<td>Synthetics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Scouring</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cotton</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Viscose</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Acetates</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Synthetics</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Wool (yarn and</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>fabric)</td>
<td>NH₄⁺</td>
<td></td>
</tr>
<tr>
<td>Wool (loose fibre)</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Bleaching</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cotton</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Linen</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Viscose</td>
<td>NH₄⁺</td>
<td></td>
</tr>
<tr>
<td>Jute</td>
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<td></td>
</tr>
<tr>
<td>Acetates</td>
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<td></td>
</tr>
<tr>
<td>Wool</td>
<td>Na⁺</td>
<td></td>
</tr>
<tr>
<td>Mercerising</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cotton</td>
<td>Na⁺NH₄⁺</td>
<td></td>
</tr>
<tr>
<td>Linen</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carbonising</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wool</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fulling</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wool</td>
<td>Na⁺</td>
<td></td>
</tr>
</tbody>
</table>

*Carboxymethyl cellulose (SB): Enzymes (A); Fats (SB); Hemicelluloses (A); Modified starches (B); Non-ionic surfactants (A); Oils (SB); Starch (B); Waxes (SB); Carboxymethyl cellulose (SB); Enzymes (A); Fats (SB); Gelatine (A); Oils (SB); Polymeric sizes (NB); Polyvinyl alcohol (A); Starch (B); Waxes (SB); Anionic surfactants (A); Cotton waxes (NB); Fats (SB); Glycerol (B); Hemicelluloses (A); Non-ionic surfactants (A); Peptic matter (A); Sizes (A); Soaps (A); Starch (A); Anionic detergents (B); Fats (SB); Non-ionic detergents (B); Oils (SB); Sizes (B); Soaps (B); Waxes (SB); Anionic surfactants (A); Anti static agents (NB); Fats (SB); Non-ionic surfactants (A); Oils (SB); Petroleum spirit (A); Sizes (B); Soaps (A); Waxes (SB); Anionic detergents (A); Glycol (SB); Mineral oils (SB); Non-ionic detergents (A); Soaps (A); Acetate (B); Anionic surfactants (A); Formate (B); Nitrogenous matter (U); Soaps (A); Suint (A); Wool grease (SB); Woll wax (SB); Oxalate (B); Alcohol sulphates (A); Anionic surfactants (A); Cresols (A); Cyclohexanol (A); Suint (A); Surfactants (A); Wool grease (SB).
### Table 3.23 (continued)

<table>
<thead>
<tr>
<th>Process/fibres</th>
<th>Substances</th>
<th>Organic (biodegradability)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dyeing</td>
<td>Inorganic</td>
<td></td>
</tr>
<tr>
<td>Cotton</td>
<td>Na⁺ Cr⁵⁺</td>
<td>CO₃²⁻ F⁻</td>
</tr>
<tr>
<td>Viscose</td>
<td>Cu²⁺</td>
<td>NO₂⁻ O₂⁻²</td>
</tr>
<tr>
<td>Linen</td>
<td>Sb¹⁺</td>
<td>S²⁻ S₂O₃²⁻</td>
</tr>
<tr>
<td></td>
<td>K⁺</td>
<td>SO₄²⁻</td>
</tr>
<tr>
<td></td>
<td>NH₄⁺</td>
<td>SO₄²⁻</td>
</tr>
<tr>
<td>Wool</td>
<td>Na⁺</td>
<td>CO₃²⁻ Cl⁻</td>
</tr>
<tr>
<td>Polyamide</td>
<td>Na⁺</td>
<td>SO₄²⁻</td>
</tr>
<tr>
<td>Acrylic</td>
<td>Na⁺</td>
<td>Cl⁻</td>
</tr>
<tr>
<td></td>
<td>Cu⁺</td>
<td>SO₄²⁻</td>
</tr>
<tr>
<td>Polyester</td>
<td>Na⁺</td>
<td>Cl⁻</td>
</tr>
<tr>
<td></td>
<td>NH₄⁺</td>
<td>Cl⁻</td>
</tr>
<tr>
<td>Fireproofing</td>
<td>Cotton</td>
<td>Ni⁺²⁺ PO₄³⁻</td>
</tr>
<tr>
<td></td>
<td>Wool</td>
<td>Na⁺ B</td>
</tr>
<tr>
<td>Moistproofing</td>
<td>Wool</td>
<td>Na⁺ K⁺ Al⁺⁺</td>
</tr>
<tr>
<td>Waterprooﬁng</td>
<td>Cotton</td>
<td>Na⁺ Cl⁻</td>
</tr>
<tr>
<td></td>
<td>Linen</td>
<td>K⁺ SO₄²⁻</td>
</tr>
<tr>
<td></td>
<td>Wool</td>
<td>Al⁺⁺</td>
</tr>
</tbody>
</table>

* B, biodegradable; A, biodegradable after acclimatisation; U, unknown; NB, non-biodegradable; SB, slowly degradable.

Chelating agents are generally used during bleaching or in the final rinses, contributing to the pollution load (Cooper, 1978; Nolan, 1972). Bleaching wastewater usually has a high solids content with low to moderate BOD levels (Table 3.21). The dissolved oxygen content of these effluents may be raised by the decomposition of hydrogen peroxide (Porter, 1990), but residual hydrogen
peroxide can cause toxicity problems in biological treatment processes (Cooper, 1978). Only light bleaching, if any, is required when processing 100% synthetic or woollen goods, and the generated wastewater is not a significant source of pollution in such cases (Nolan, 1972).

**Mercerising**
Mercerisation is performed almost exclusively on pure cotton fabrics, which are treated by a concentrated caustic bath and a final acid wash to neutralise them. Its purpose is to give lustre and also to increase dye affinity and tensile strength. Mercerisation wastewaters have low BOD and total solids levels but are highly alkaline prior to neutralisation (Cooper, 1978; Nolan, 1972). The low BOD content arises from surfactants and penetrating agents used as auxiliary chemicals (Table 3.21).

**Carbonising**
Carbonisation is performed on woollen items to remove traces of vegetable matter. The process can be carried out either in conjunction with raw scouring or at the fabric processing stage, depending on the level of impurities and the end use of the wool (Cooper, 1978; OECD, 1981). Carbonising consists of soaking the material in dilute sulphuric acid followed by neutralisation with sodium carbonate. The material is then dried and the brittle cellulosic matter mechanically removed. The generally low levels of organic materials in carbonisation effluents are due to vegetable matter, whilst the acid treatment yields high levels of dissolved solids. Performing carbonisation in conjunction with raw wool scouring leads to a reduction in the total pollution load of the scouring waste stream (OECD, 1981).

**Fulling**
Fulling stabilises woollen fabrics and gives them a thicker and more compact appearance. It is carried out with soda ash or sulphuric acid in the presence of detergents, sequestering agents, metallic catalysts and hydrogen peroxide in conjunction with mechanical agitation (Nolan, 1972; OECD, 1981). The fulling solution is then drained and the treated product extensively washed to remove the remaining chemicals. Fulling wastes in combination with effluents generated by subsequent washing operations present, after raw wool scouring, the largest source of BOD in wool processing wastewaters (Cooper, 1978; Nolan, 1972). Most of the BOD arises from soap, detergents and lubricants and oils added to the wool during the production process.

**Dyeing**
Dyeing is carried out to add colour to fabrics or yarn. Identification of generic types of dyeing wastewaters is complicated by the diversity of both the dye chemistry and the operational modes of the dyeing process itself. Although rarely toxic, these wastewaters demand special consideration since they are arguably the most problematic of all textile wastewaters, for a number of reasons:
they are produced in large volumes (around 100–150 kg\(^{-1}\) textile product for the average dyeing and rinsing operation),

- they are not readily biodegradable, such that conventional municipal wastewater treatment plant will generally remove only around 20–30% of colour associated with synthetic dyes, and

- they require removal to very low levels prior to discharge if consents based on colour are in place.

Dyes are generally small molecules comprising two key components: the chromophores, responsible for the colour, and the auxochromes, which can not only supplement the chromophore but also render the molecule soluble in water and give enhanced affinity toward the fibres (Trotman, 1984). A large number of dyes are reported in specialised literature (Colour Index, 1987). These can be classified both by their chemical structure or their application to the fibre type (Table 3.24). Dyes may also be classified on the basis of their solubility: soluble dyes include acid, mordant, metal complex, direct, basic and reactive dyes; and insoluble dyes include azoic, sulphur, vat and disperse dyes. An alternative dye classification that refers to colour removal technologies (Treffry-Goatley and Buckley, 1991) places the various classes of dyes (with respect to their application) into three groups depending on their state in solution and on the type of charge the dye acquires. Each group can be associated with potential colour-removal methods (Table 3.25).

Complex chemical and/or physical mechanisms govern the adsorption and retention of dyes by fibres. The adsorptive strength, levelling and retention are controlled by several factors such as time, temperature, pH, and auxiliary chemicals (Nunn, 1979; Trotman, 1984; Preston, 1986; Shore, 1990). A large range of substances other than dyes, auxiliary chemicals used in the dyeing process, can be found in a dye effluent at any one time. The effluent composition and colour is further complicated by the fact that both dye fixation rates (Table 3.26) and liquor ratios (the volume of dye solution per weight of goods) vary, and different dye classes may be used for a single dyeing operation (Shore, 1990; Horning, 1978). Moreover, continuous operation yields smaller volumes of more concentrated dyewaste than batch operation, equating to typically a four-fold factorial difference with respect to dye concentration and a 2.5-fold difference in volume (Glover and Hill, 1993), some typical batch process effluent data being given in Table 3.27.

**Chemical finishing**

Chemical finishing processes include processes designed to change the optical, tactile, mechanical strength or dirt-releasing properties of the textile. Optical finishes can either brighten or deluster the textile (NCDNER, 1995; OECD, 1981). Softeners and abrasion-resistant finishes are added to improve the feel or increase the ability of the textile to resist abrasion and tearing. Absorbent and soil release finishes alter the surface tension and other properties to increase water absorbency or improve soil release. Physical stabilisation and crease-resistant finishes, which may include formaldehyde-based resin finishes,
<table>
<thead>
<tr>
<th>Chemical class</th>
<th>Colour index number</th>
<th>Fibre type/dye class</th>
<th>Cellulosic</th>
<th>Protein</th>
<th>Cellulose ester</th>
<th>Polyamide</th>
<th>Polyester</th>
<th>Polyacrylic</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Direct</td>
<td>Sulphur</td>
<td>Azoic</td>
<td>Vat</td>
<td>Reactive</td>
<td>Acid</td>
</tr>
<tr>
<td>Nitroso</td>
<td>10000-10299</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Nitro</td>
<td>10300-10999</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Azo</td>
<td>11000-36999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Azoic</td>
<td>37000-39999</td>
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<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Stilbene</td>
<td>40000-40799</td>
<td></td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Carotenoid</td>
<td>40800-40999</td>
<td></td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Triarylmethane</td>
<td>42000-44999</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Xanthenic</td>
<td>45000-45999</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Acridine</td>
<td>46000-46999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Quinoline</td>
<td>47000-47999</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Methine</td>
<td>48000-48999</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Thiazole</td>
<td>49000-49399</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Oxazine</td>
<td>50000-50999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Thiazine</td>
<td>51000-51999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Sulphur</td>
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<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Anthraquinone</td>
<td>53000-54999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Indigoid</td>
<td>58000-72999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Phthalocyanine</td>
<td>74000-74999</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

*Does not include all chemical classes.*
Table 3.25  Dye classification with respect to colour removal (Treffry-Goatley and Buckley, 1991)

<table>
<thead>
<tr>
<th>Classification</th>
<th>Dye class</th>
<th>Charge/solution state</th>
<th>Technology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I</td>
<td>Disperse</td>
<td>Negatively charged</td>
<td>Coagulation</td>
</tr>
<tr>
<td></td>
<td>Azoic</td>
<td>Colloidal</td>
<td>Membrane Oxidation</td>
</tr>
<tr>
<td></td>
<td>Vat</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sulphur</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group A</td>
<td>Acid</td>
<td>Anionic</td>
<td>Adsorption</td>
</tr>
<tr>
<td></td>
<td>Reactive</td>
<td>Soluble</td>
<td>Ion exchange</td>
</tr>
<tr>
<td></td>
<td>Direct</td>
<td></td>
<td>Membrane Oxidation</td>
</tr>
<tr>
<td></td>
<td>Mordant</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Metal complex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group C</td>
<td>Basic</td>
<td>Cationic</td>
<td>Adsorption</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Soluble</td>
<td>Ion exchange</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Membrane Oxidation</td>
</tr>
</tbody>
</table>

Table 3.26  Percentage unfixed dye for different dye types and applications (ETBP, 1997)

<table>
<thead>
<tr>
<th>Fibre</th>
<th>Dye type</th>
<th>Unfixed dye, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wool and nylon</td>
<td>Acid/reactive dyes for wool</td>
<td>7–20</td>
</tr>
<tr>
<td></td>
<td>Pre-metallised</td>
<td>2–7</td>
</tr>
<tr>
<td></td>
<td>Chromic</td>
<td>1–2</td>
</tr>
<tr>
<td>Cotton and viscose</td>
<td>Azoic</td>
<td>5–10</td>
</tr>
<tr>
<td></td>
<td>Reactive</td>
<td>20–50</td>
</tr>
<tr>
<td></td>
<td>Direct</td>
<td>5–20</td>
</tr>
<tr>
<td></td>
<td>Pigment</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Vat</td>
<td>5–20</td>
</tr>
<tr>
<td></td>
<td>Sulphur</td>
<td>30–40</td>
</tr>
<tr>
<td>Polyester</td>
<td>Disperse</td>
<td>8–20</td>
</tr>
<tr>
<td>Acrylic</td>
<td>Modified</td>
<td>2–3</td>
</tr>
</tbody>
</table>

stabilise cellulosic fibres to laundering and shrinkage, imparting permanent press properties to fabrics. Finishing processes generally involve impregnation of the fabric using a padding and mangle technique followed by a fixation step by heat. Subsequent washing may be carried out to remove residual chemicals. Whilst low in volume, the effluents from these finishing operations are extremely variable in composition and can contain toxic organic substances such as pentachlorophenols and ethylchlorophosphates (Table 3.23).

3.3.3 Process water quality requirements

It is generally the case that, in common with many industrial activities, textile processing makes use of mains water whose quality is therefore stipulated by statutory regulations for drinking water. However, water is also provided by abstraction from ground, lake and river water – options made more attractive by
Table 3.27 Characteristics of dyeing effluents (Horning, 1978)

<table>
<thead>
<tr>
<th>Dye</th>
<th>Fibre</th>
<th>Colour (ADMI) (mg/l)</th>
<th>BOD (mg/l)</th>
<th>TOC (mg/l)</th>
<th>TSS (mg/l)</th>
<th>TDS (mg/l)</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acid</td>
<td>Polyamide</td>
<td>4000</td>
<td>240</td>
<td>315</td>
<td>14</td>
<td>2028</td>
<td>5.1</td>
</tr>
<tr>
<td>Mordant</td>
<td>Wool</td>
<td>3200</td>
<td>135</td>
<td>210</td>
<td>9</td>
<td>1086</td>
<td>4.0</td>
</tr>
<tr>
<td>1:2 Metal complex</td>
<td>Polyamide</td>
<td>370</td>
<td>570</td>
<td>400</td>
<td>5</td>
<td>3945</td>
<td>6.8</td>
</tr>
<tr>
<td>Basic</td>
<td>Acrylic</td>
<td>5600</td>
<td>210</td>
<td>255</td>
<td>13</td>
<td>1469</td>
<td>4.5</td>
</tr>
<tr>
<td>Basic</td>
<td>Polyester</td>
<td>1300</td>
<td>1470</td>
<td>1120</td>
<td>4</td>
<td>1360</td>
<td>5.0</td>
</tr>
<tr>
<td>Direct</td>
<td>Viscose</td>
<td>12500</td>
<td>15</td>
<td>140</td>
<td>26</td>
<td>2669</td>
<td>6.6</td>
</tr>
<tr>
<td>Direct (cooper treated)</td>
<td>Cotton</td>
<td>525</td>
<td>87</td>
<td>135</td>
<td>41</td>
<td>2763</td>
<td>5.0</td>
</tr>
<tr>
<td>Reactive (batch)</td>
<td>Cotton</td>
<td>3890</td>
<td>0</td>
<td>150</td>
<td>32</td>
<td>12500</td>
<td>11.2</td>
</tr>
<tr>
<td>Reactive (continuous)</td>
<td>Cotton</td>
<td>1390</td>
<td>102</td>
<td>230</td>
<td>9</td>
<td>691</td>
<td>9.1</td>
</tr>
<tr>
<td>Azoic</td>
<td>Cotton</td>
<td>2415</td>
<td>200</td>
<td>170</td>
<td>387</td>
<td>10900</td>
<td>9.3</td>
</tr>
<tr>
<td>Sulphur (continuous)</td>
<td>Cotton</td>
<td>450</td>
<td>990</td>
<td>400</td>
<td>34</td>
<td>2000</td>
<td>3.7</td>
</tr>
<tr>
<td>Vat</td>
<td>Cotton</td>
<td>1910</td>
<td>294</td>
<td>265</td>
<td>41</td>
<td>3945</td>
<td>11.8</td>
</tr>
<tr>
<td>Disperse (high temp.)</td>
<td>Polyester</td>
<td>1245</td>
<td>198</td>
<td>360</td>
<td>76</td>
<td>1700</td>
<td>10.2</td>
</tr>
<tr>
<td>Disperse (Atmosf. dyeing)</td>
<td>Polyester</td>
<td>315</td>
<td>234</td>
<td>300</td>
<td>39</td>
<td>914</td>
<td>7.8</td>
</tr>
<tr>
<td>Disperse</td>
<td>Polyester (carpet)</td>
<td>215</td>
<td>159</td>
<td>240</td>
<td>101</td>
<td>771</td>
<td>7.1</td>
</tr>
<tr>
<td>Disperse</td>
<td>Polyamide (carpet)</td>
<td>100</td>
<td>78</td>
<td>130</td>
<td>14</td>
<td>396</td>
<td>8.3</td>
</tr>
<tr>
<td>Disperse/acid/basic (continuous)</td>
<td>Polyamide (carpet)</td>
<td>130</td>
<td>160</td>
<td>49</td>
<td>258</td>
<td>6.5</td>
<td></td>
</tr>
<tr>
<td>Disperse/acid/basic (batch)</td>
<td>Polyamide (carpet)</td>
<td>210</td>
<td>42</td>
<td>130</td>
<td>8</td>
<td>450</td>
<td>6.7</td>
</tr>
<tr>
<td>Disperse/vat (continuous)</td>
<td>Cotton/polyester</td>
<td>365</td>
<td>360</td>
<td>350</td>
<td>9</td>
<td>691</td>
<td>9.1</td>
</tr>
</tbody>
</table>

The spiralling costs of mains water. Guide values for key water quality determinants are included in Table 3.28 (Mattioli et al., 2002), which includes process water quality data for three operational textile finishing plants in the Como and Prato textile production areas of Italy. It is evident from these guide values that the water quality requirement is less rigorous than that demanded for drinking water by European Union legislation, which additionally has to be bacteriologically safe, such that tertiary-treated wastewater is apparently suitable for reuse (Section 3.1.7).

3.3.4 Legislation

USA

The most important environmental regulation affecting the textile industry in the USA is the Clean Water Act. In 1982 the US Environmental Protection Agency (EPA) promulgated effluent guidelines for the textile manufacturing point source category. The Textile Mills Point Source Category effluent guidelines are listed under 40 CFR (Code of Federal Regulations) Part 410. Part 410 is divided into nine subparts, each applying to a different textile manufacturing subcategory as outlined below. Each subpart contains effluent limitations, new source performance standards (NSPS), and pre-treatment standards. The definition of Subpart D for the woven fabric includes desizing.
**Table 3.28** Average process water quality for textile finishing (Mattioli et al., 2002)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Lake</th>
<th>Ground</th>
<th>Reclaimed*</th>
<th>Guide value</th>
</tr>
</thead>
<tbody>
<tr>
<td>TSS, mg/l</td>
<td>&lt;1</td>
<td>10</td>
<td>22</td>
<td>10</td>
</tr>
<tr>
<td>COD, mg/l</td>
<td>&lt;10</td>
<td>30</td>
<td>53</td>
<td>30</td>
</tr>
<tr>
<td>UV abs. 420 nm</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.02</td>
<td>0.01</td>
</tr>
<tr>
<td>pH</td>
<td>7.8</td>
<td>7.5</td>
<td>7.7</td>
<td>7–8</td>
</tr>
<tr>
<td>Conductivity, μS/cm</td>
<td>290</td>
<td>200</td>
<td>1600</td>
<td>1800</td>
</tr>
</tbody>
</table>

*Reclaimed from tertiary-treated wastewater (Section 3.1.7).

bleaching, mercerising, dyeing, printing, resin treatment water proofing, flame proofing, soil repellency application, and special finish application.

Effluent limitations representing the degree of effluent reduction attainable by using either best practicable control technologies (BPT), or best available technologies (BAT) are given for all subcategories. BPTs are used for discharges from existing point sources to control conventional and non-conventional pollutants as well as some priority pollutants. BATs are used to control priority pollutants and non-conventional pollutants when directly discharged into environmental waters. BPT limits for BOD, COD, TSS, sulphide, phenol, total chromium, and pH are set for every category (Subpart), with the exception of Subpart C (Low Water Use Processing Subcategory). Each subpart, with the exception again of Subpart C, also has BAT limits for COD, sulphide, phenols, and total chromium. New source performance standards (NSPS) for BOD, COD, TSS, sulphide, phenols, total chromium, and pH are set for each subcategory. However, for the woven fabric finishing subcategory (Subpart D), the NSPSs are divided into process-specific standards that differ according to the manufacturing operation. The NSPS levels (Table 3.29) are based on mass emission rather than concentration.

The USEPA has also recently developed values for factors related to effluent generation load factors (i.e. mass flow of pollutant per based on unit production) to enable estimation of the effluent load from different textile mills (Table 3.30). In the absence of actual data, these estimates provide some guideline to estimate the extent of environmental impact. Existing textile mills, on the other hand, are able to use this data for benchmarking.

**Europe**

European legislation is less prescriptive than that of the USEPA, but nonetheless is based on the same principle of BAT. The key European directive is 96/61 Integrated Pollution Prevention and Control, which requires both industrial plant operators and regulators to take an integrated, holistic view of the pollution and resource demand potential of the installation. Central to this approach is appropriate preventative measures against pollution, specifically through the application of the BAT principle to improve environmental performance (Dulio, 2001). It is then left to the individual member states to define BATs, but these must obviously focus on waste minimisation and, ultimately, closed-loop options for industrial water use.
Table 3.29  NSPSs for liquid effluents from cotton woven fabric finishing: 30 day average (1 day maximum) in kg/MT of product (EPA, 1997)

<table>
<thead>
<tr>
<th>Process</th>
<th>BOD</th>
<th>COD</th>
<th>TSS</th>
<th>Sulphide</th>
<th>Phenol</th>
<th>Chromium</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simple</td>
<td>1.7 (3.3)</td>
<td>26.9 (41.7)</td>
<td>3.9 (8.8)</td>
<td>0.1 (0.2)</td>
<td>0.05 (0.1)</td>
<td>0.05 (0.1)</td>
<td>(6–9)</td>
</tr>
<tr>
<td>Complex</td>
<td>1.9 (3.7)</td>
<td>44.2 (68.7)</td>
<td>6.4 (14.4)</td>
<td>0.1 (0.2)</td>
<td>0.05 (0.1)</td>
<td>0.05 (0.1)</td>
<td>(6–9)</td>
</tr>
<tr>
<td>Desizing</td>
<td>2.8 (5.5)</td>
<td>38.3 (59.5)</td>
<td>6.9 (15.6)</td>
<td>0.1 (0.2)</td>
<td>0.05 (0.1)</td>
<td>0.05 (0.1)</td>
<td>(6–9)</td>
</tr>
</tbody>
</table>

Table 3.30  Pollutant load in kg m⁻³, median values, from woven fabric finishing (UNEP, 1992)

<table>
<thead>
<tr>
<th>Sub-category</th>
<th>BOD</th>
<th>COD</th>
<th>TSS</th>
<th>Oil and grease</th>
<th>Phenol</th>
<th>Cr</th>
<th>Sulphide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simple processing</td>
<td>22.6</td>
<td>92.4</td>
<td>8.0</td>
<td>9.1</td>
<td>8.2</td>
<td>4.3</td>
<td>7.6</td>
</tr>
<tr>
<td>Complex processing</td>
<td>32.7</td>
<td>110.6</td>
<td>9.6</td>
<td>3.8</td>
<td>7.7</td>
<td>2.6</td>
<td>12.5</td>
</tr>
<tr>
<td>Complex processing +</td>
<td>45.1</td>
<td>122.6</td>
<td>14.8</td>
<td>4.1</td>
<td>13.1</td>
<td>20.9</td>
<td>N</td>
</tr>
<tr>
<td>desizing</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

aN, sufficient data not available to report.

There exist examples of regulatory bodies imposing discharge consents relating to textile processing activities. One of these concerns the Wigston sewage treatment works in the Severn Trent region of the UK, to which a number of dyehouses were discharging coloured effluents. This resulted in the sewage treatment works, which was of conventional design, discharging coloured effluent into the River Sence. The Environment Agency imposed colour consent limits on the works, which Severn Trent Water, the sewerage operator, passed on to the five dyehouses discharging to the sewage treatment works. This dramatically influenced the economics of effluent recycling, as is explained in the case study in Section 5.7.

3.3.5 Conventional treatment

Conventional technologies for textile wastewater treatment mainly comprise biological treatment, precipitation, coagulation/flocculation, flotation, oxidation and adsorption (Table 3.31), and have recently been reviewed (Vandevivere et al., 1998; Slokar and Le Marechal, 1998; Rott and Minke, 1999; Delee et al., 2002). Since the bulk of the COD for many textile wastewaters is biodegradable, most textile effluents are discharged to sewer, sometimes following rudimentary aerobic biological treatment. For high COD loads, such as those arising from desizing of starch or wool washing and scouring, anaerobic pretreatment can be employed (Sacks and Buckley, 1999; Shaw et al., 2002; Delée et al., 2002), although this appears not to have reached commercial scale. Coagulation, often followed by sand filtration, is also extensively used.

The problems imposed by dyewaste, however, are particularly vexing. Biodegradability studies have indicated the feasibility of reducing the BOD and
Table 3.31 Technologies for textile effluent treatment (adapted from Vandevivere et al., 1998)

<table>
<thead>
<tr>
<th>Process</th>
<th>Stage</th>
<th>Status</th>
<th>Performance</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biodegradation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Activated sludge</td>
<td>Main treatment</td>
<td>Widely used</td>
<td>Bulk COD removal: partial nitrification</td>
<td>High residual COD, ammonia, colour</td>
</tr>
<tr>
<td>Sequential anaerobic-aerobic</td>
<td>Main treatment</td>
<td>Few reports of full-scale use</td>
<td>Better COD and colour removal than AS alone</td>
<td>High residual COD, colour</td>
</tr>
<tr>
<td>Fixed bed</td>
<td>Main treatment</td>
<td>Pilot trials in China</td>
<td>Better COD and colour removal than AS</td>
<td>Some residual colour</td>
</tr>
<tr>
<td>Fungi/H$_2$O$_2$</td>
<td>Main treatment</td>
<td>Bench-scale</td>
<td>Full decolourisation</td>
<td></td>
</tr>
<tr>
<td>Physicochemical treatment</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coagulation-flocculation</td>
<td>Pre-, main or post-treatment</td>
<td>Extensive use</td>
<td>Almost full decolourisation; water reuse</td>
<td>Unreliable performance: sludge disposal</td>
</tr>
<tr>
<td>Adsorption</td>
<td>Pre- or post-treatment</td>
<td>Bench- to full-scale depending on adsorbent</td>
<td>Water reuse with newer adsorbents</td>
<td>Sludge disposal or adsorbent regeneration</td>
</tr>
<tr>
<td>Membrane filtration</td>
<td>Main or post-treatment</td>
<td>Extensive use in South Africa</td>
<td>Reliable performance; water reuse</td>
<td>Concentrate management</td>
</tr>
<tr>
<td>Oxidation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ozonation</td>
<td>Post-treatment</td>
<td>Full-scale</td>
<td>Full decolourisation; water reuse</td>
<td>Expensive: aldehydes formed</td>
</tr>
<tr>
<td>Fenton's reagent</td>
<td>Pre-treatment</td>
<td>Several full-scale plant in South Africa</td>
<td>Full decolourisation</td>
<td></td>
</tr>
<tr>
<td>Photocatalysis</td>
<td>Post-treatment</td>
<td>Pilot-scale</td>
<td>Near complete colour removal</td>
<td>For final polishing only</td>
</tr>
<tr>
<td>Electrolysis</td>
<td>Pre-treatment</td>
<td>Pilot-scale</td>
<td>Full decolourisation</td>
<td>Foaming: limited electrode life</td>
</tr>
</tbody>
</table>
COD or TOC of dyeing wastewaters by biological treatment (Horning, 1978; Pagga and Brown, 1986; Willmot et al., 1998; Lorenço et al., 2001). Biological colour removal is low, however, unless extended retention times (2–3 days) are employed, since most dye molecules are not biodegradable. The partial treatment of dyes by conventional biological processes is attributed to precipitation (in the case of sparingly soluble dyes) or adsorption on the sludge (Porter and Snider, 1976; Weeter and Hodgson, 1975). Reactive and acid dyes, on the other hand, are not adsorbed on biological sludge to any great degree (Hitz et al., 1978; Slokar and Le Marechal, 1998). The problem of colour from reactive dyes in dyeing effluents is further aggravated by their high commercial popularity and generally low fixation rates (as low as 50% for cotton and viscose fibres) relative to other dye classes. Thus, even though they form only around 2% of the COD in dyewaste, they present the most intractable of problems.

Several physicochemical colour removal methods have been employed at full-scale over the years (Table 3.31). Many studies have focused on ozonation (Lin and Lin, 1993; Perkowski et al., 1996; Matsui, 1996) and advanced oxidation (Solozhenkho et al., 1995; Lin and Peng, 1996) which reliably decolourise a wide range of dyewastes, although there has also been some notable success using the proprietary zeolite-based adsorbent Macrosorb. Having said this, all treatments present particular disadvantages. Advanced oxidation is high in capital and/or operating costs and may create mutagenic products. Coagulation creates significant quantities of sludge, as does powdered activated carbon. Adsorption and ion exchange are generally low in capacity, necessitating large quantities of materials. Also, and in common with membrane processes, they produce a concentrated regenerant product or else, in the case of granular activated carbon, demand an expensive regeneration process. Finally, no single process is universally effective against all dye chemicals – although reverse osmosis is certainly the most effective decolouring and desalting process against the most diverse range of dyewastes, and has been successfully employed for recycling (Buckley, 1992). Crucially, only dense membrane processes offer the opportunity to remove dissolved solids (Section 3.1.7), which would otherwise accumulate for any chemical treatment process.

### 3.3.6 Demand management

Given the exigencies of textile wastewater treatment, and of dyewaste in particular, it is unsurprising that much effort in cost and environmental impact reduction within the industry has been directed at reducing water and chemicals consumption. There are a large number of documented examples of cost savings in textile fabrication facilities through basic low-cost process modification and effective, less profligate water use. Simple reuse of process water from dyeing, rinsing and bleaching operations without further treatment, as proscribed through water auditing techniques such as pinch analysis (Section 4.2), can provide substantial cost benefits (Table 3.32). There are many other examples of cost savings made through substitution of process chemicals for ones less environmentally onerous (such as starch with PVA/acylates and soaps with
Table 3.32 Process modification cost benefit examples (EPA, 1997; ETPB, 1997)

<table>
<thead>
<tr>
<th>Company</th>
<th>Action</th>
<th>Cost benefit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lancashire medical textile company</td>
<td>Elimination of two wash cycles from bleaching process</td>
<td>£1700 reduction in effluent costs</td>
</tr>
<tr>
<td>American Corporation</td>
<td>Extension of length of dyeing time by 15 minutes to improve exhaustion</td>
<td>60% decrease in BOD and COD; 20% drop in FOG; 98% drop in NH$_3$-N; £35 000 decrease p.a.</td>
</tr>
<tr>
<td>Adams-Millis Co.</td>
<td>Reuse of dyebath for nylon hosiery in rotary drum dyeing machines</td>
<td>35% decrease in water use ($0.04/kg product)</td>
</tr>
<tr>
<td>Bigelow Carpets</td>
<td>Cycling of dyebath water between pairs of dyeing machines to allow reuse over 20 cycles</td>
<td>£60 000 p.a., plus reduction in effluent BOD and colour</td>
</tr>
<tr>
<td>A yarn finishing company</td>
<td>Reuse of rinsewater three times following mercerising, with evaporation to recover caustic Na$_2$CO$_3$</td>
<td>Reduction of 80% TSS, 55% COD and 70% in Na$_2$CO$_3$ in effluent (and so HCl for neutralisation). £189 000 p.a. (1 year payback) 50% reduction in effluent COD £2700 p.a.</td>
</tr>
<tr>
<td>North Carolina facility</td>
<td>Bleach bath reuse</td>
<td>£10 700 p.a. in effluent disposal and water consumption reduction</td>
</tr>
<tr>
<td>Lancashire medical textile company</td>
<td>Recycling of last rinse from beam-and-winch bleaching operation</td>
<td></td>
</tr>
<tr>
<td>Wigan dyehouse</td>
<td>Reuse of effluent from bleaching for scouring wash</td>
<td></td>
</tr>
</tbody>
</table>

synthetic detergents, in both cases to reduce effluent BOD levels), and reuse of other process waters, such as coolant (ETPB, 1997). None of these measures, however, can substantially reduce wastewater discharge to the extent possible through effluent recycling.

3.3.7 Reuse practice and opportunities

Reclamation and reuse
Given the capital intensive nature of ozonation, which represents arguably the most reliable and established decolourisation process, it is perhaps unsurprising that there exist examples of centralised wastewater treatment facilities based on ozonation (Churchley, 1994; Mattioli et al., 2002). For one region, the Como province of northern Italy, there exist two sewage treatment works (Bulgarograsso STW and Fino Mornasco STW) designed to treat wastewater of which around 50% originates from industrial activity and, of this, over 90% from textile processing operations. Both plants, which have a combined capacity of 100 000 m$^3$ per day, are based on conventional primary and secondary treatment with ozonation tertiary treatment following pretreatment either by sand filtration or coagulation/flocculation with lamella settling. Further polishing of the effluent from these works using fixed-film granular activated carbon, i.e. biologically activated carbon or BAC, produced water of a quality found to be suitable for silk degumming and dyeing (Mattioli et al., 2002). Although not extensive, the trial appears to demonstrate the feasibility of recycling of dyewaste that has undergone ozonation tertiary treatment.
Recycling

The textile processes providing the most propitious conditions for application of membranes are fabric coating, desizing, scouring and dyeing. They have been successfully employed at full scale in all these areas for the last 20 years in South Africa, and more recently in other countries.

For two of these operations both the water and the concentrate are reusable. Fabric coating makes use of latex, a relatively expensive chemical, which arises in low concentrations in the effluent as an emulsion, with droplet sizes down to 0.05 μm. The use of ultrafiltration for latex and water recovery allows substantial recovery of the emulsified latex (Groves et al., 1978). The main limitation appears to be from the stability of the latex. Coagulation of the latex on the membrane produces a deposit that can only be removed using appropriate organic solvents and is then no longer recoverable. On the other hand, latex stability can be improved through dosing with appropriate surfactants (Cheryan, 1998). Recovery of synthetic size (polyvinyl alcohol and polyacrylate) is also possible using UF (Porter, 1998), with original reports of successful pilot-scale trials dating back to 1980 (Groves and Buckley, 1980). In this instance the elevated effluent temperature (75°C) both increases the permeate flux and also increases the cost benefit through recovering energy as heat.

Wool scouring produces a high-BOD effluent (Table 3.20) containing detergents, suspended solids, fats, oils and grease (FOG) which can be treated biologically and/or by dissolved air flotation. The use of ultrafiltration, operating at 8–10 bar, permits substantial concentration of the major pollutants in the retentate stream. The concentration factor is typically ~7, producing a retentate very high in organics (10–35 wt% COD). The longest established wool scouring UF plant in Europe is in Norway (Bilstad et al., 1994). This UF plant has been operational since 1989 and achieves >80% COD removal, nonetheless leaving a filtrate still too high in COD for recycling. An interesting recent development is the emergence of membrane bioreactors for this duty. A xenon hollow fibre submerged MBR has been installed for treating a 13 m³ h⁻¹ flow of 3500–5000 mg l⁻¹ COD wool scouring wastewater in the UK (Bennett, 2000). Polishing of the effluent by a two-stage reverse osmosis concentrator process, which includes interstage pumping, which is designed to attain an overall recovery of 85% (75% from the first stage and 50% from the second). No performance data has been reported for this system, however.

Reviews of the use of membranes for dyewastewater treatment have been presented by a number of authors (Buckley, 1992; Cooper, 1993; Diaper et al., 1996). Although not all membrane applications relate specifically to recycling, it is apparent from the performance attainable in terms of rejection and product water quality (Tables 3.33 and 3.34) that recycling of the permeate is possible in many cases. At least two of the more recent studies (Marcucci et al., 2001; Sójka-Ledakowicza et al., 1998) listed in Table 3.34 have incorporated testing of the recovered water for dyeing. In both cases, it was concluded that the more selective reverse osmosis process was required for a reusable water product.

Dyewaste is characteristically high in dissolved solids and low in suspended solids, with colour arising from exhausted dye (Table 3.20). It is of interest to
<table>
<thead>
<tr>
<th>Reference</th>
<th>Dye type</th>
<th>Removal process used</th>
<th>Scale, flow (m³/day) or membrane area (m²)</th>
<th>Permeate flux (l/m²h)</th>
<th>Dyestuff reused</th>
<th>% colour rejection</th>
<th>Permeate reused?</th>
<th>% salt rejection</th>
<th>% water recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buckley (1992)</td>
<td>reactive soluble and colloidal</td>
<td>NF</td>
<td>0.56 m² membrane area</td>
<td>33</td>
<td>NO</td>
<td>99</td>
<td>NO</td>
<td>80</td>
<td>85</td>
</tr>
<tr>
<td>Buckley (1992)</td>
<td>disperse soluble and colloidal</td>
<td>dynamic-formed UF</td>
<td>256 m² membrane area</td>
<td>No</td>
<td>-95</td>
<td>Yes</td>
<td>81</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buckley (1991)</td>
<td>reactive soluble and colloidal</td>
<td>MF</td>
<td>40 m³/day</td>
<td>50</td>
<td>No</td>
<td>81</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gaeta and Vedele (1991)</td>
<td>reactive soluble and colloidal</td>
<td>MF</td>
<td>15 m³/day</td>
<td>15</td>
<td>No</td>
<td>81</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calabro et al. (1991)</td>
<td>reactive soluble and colloidal</td>
<td>RO</td>
<td>lab scale</td>
<td>variable</td>
<td>No</td>
<td>91-94</td>
<td>Yes</td>
<td>98</td>
<td>85 95</td>
</tr>
<tr>
<td>Majewska-Nowak et al. (1984)</td>
<td>reactive soluble and colloidal</td>
<td>NF</td>
<td>15.9 x 10⁻⁴ m² membrane area</td>
<td>2.1 8.3</td>
<td>membrane area</td>
<td>95-100°</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Majewska-Nowak et al. (1989)</td>
<td>reactive soluble and colloidal</td>
<td>UF, polysulphone membranes</td>
<td>81.2 x 10⁻⁴ m² membrane area</td>
<td>33.3-41 7</td>
<td>membrane area</td>
<td>97b</td>
<td>10-20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porter (1990)</td>
<td>reactive soluble and colloidal</td>
<td>UF</td>
<td>150 m³/day</td>
<td>33</td>
<td>No</td>
<td>97</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Townsend et al. (1992)</td>
<td>reactive soluble and colloidal</td>
<td>UF</td>
<td>40 m³/day</td>
<td>50</td>
<td>No</td>
<td>97</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Treffry-Goatley and Buckley (1991)</td>
<td>reactive soluble and colloidal</td>
<td>UF, polysulphone membranes</td>
<td>15.9 x 10⁻⁴ m² membrane area</td>
<td>5.8</td>
<td>membrane area</td>
<td>90-100°</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Majewska-Nowak et al. (1984)</td>
<td>reactive soluble and colloidal</td>
<td>NF, ZrO-PAA on sintered stainless steel</td>
<td>139 m² membrane area</td>
<td>139</td>
<td>Yes</td>
<td>97</td>
<td>(partial)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porter and Goodman (1984)</td>
<td>reactive soluble and colloidal</td>
<td>NF, ZrO-PAA on sintered stainless steel</td>
<td>139</td>
<td>83</td>
<td>lab</td>
<td>75-80</td>
<td>(Table continued on next page)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Note:* The table continues on the next page with additional rows and data.
Table 3.33 (continued)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Dye type</th>
<th>Scale, flow (m3/day)</th>
<th>Permeate dye</th>
<th>% colour</th>
<th>Permeate salt</th>
<th>% reused</th>
<th>Permeate water</th>
<th>% recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calabro * et al. (1990)</td>
<td>Bayer Levax: red, blue</td>
<td>RO</td>
<td>5 m² membrane area</td>
<td>5.2 x 10^-4 m²</td>
<td>81.2 x 3.75-5.0</td>
<td>97.80</td>
<td>12.9-15.4</td>
<td>82.2-15.3</td>
</tr>
<tr>
<td>Watters et al. (1991)</td>
<td>Analogues: direct green, UF: polysulphone</td>
<td>UF</td>
<td>0.11 m² membrane area</td>
<td>27.5-370</td>
<td>34-93</td>
<td>No</td>
<td>85-95</td>
<td></td>
</tr>
</tbody>
</table>

* For dyes with molecular weight > 900.  
* For dyes with molecular weight > 780.  
* For dyes with molecular weight > 739.
Table 3.34  Recent publications on membranes for dyestuff removal

<table>
<thead>
<tr>
<th>Reference</th>
<th>Dye or matrix</th>
<th>Removal process</th>
<th>Membrane area (m²)</th>
<th>Permeate flux (l/m²h)</th>
<th>Colour rejection (or COD)</th>
<th>Permeate reused?</th>
<th>% salt rejection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akbari et al. (2002)</td>
<td>Acid, direct, disperse, reactive</td>
<td>NF (bespoke)</td>
<td>0.00132</td>
<td>6–18 at 1–4 bar</td>
<td>87–100%</td>
<td>No</td>
<td>~10</td>
</tr>
<tr>
<td>Bes-Pia et al. (2002)</td>
<td>Real effluent</td>
<td>Pre-coag. UF, NF (NF70, Film-Tec)</td>
<td>0.009</td>
<td>60 at 1.5 bar, UF; 22, NF</td>
<td>&gt; 97% (COD)</td>
<td>No</td>
<td>85%</td>
</tr>
<tr>
<td>Yen et al. (2002)</td>
<td>Real effluent</td>
<td>RO: Film-Tec TW30-1812-50</td>
<td>0.0017</td>
<td>40–60 at 3.4–14 bar, 45°C</td>
<td>72–98% at 3.4–14 bar, 45°C</td>
<td>No</td>
<td>85–98% at 3.4–14 bar and 45–15°C</td>
</tr>
<tr>
<td>Voigt et al. (2001)</td>
<td>Real effluent</td>
<td>TiO₂ NF, MWCO 450 Da</td>
<td>5.1</td>
<td>39–215 at 6–25 bar; 12–60 at 2.5–5 m/s, 15 bar</td>
<td>70–100%</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>Marlucci et al. (2001); Ciardelli et al. (2001)</td>
<td>Real effluent</td>
<td>PVDF UF, MWCO 70k Da; Osmonics NF, Dl.4040F SW, 130–300 Da, Film-Tec RO</td>
<td>Single element</td>
<td>–</td>
<td>90% by RO, 0% by UF</td>
<td>Yes (for RO – NF insufficient for reuse)</td>
<td>40% (NF)</td>
</tr>
<tr>
<td>Sójka-Ledakovicz et al. (1998)</td>
<td>Analogue and real effluent</td>
<td>Desalination Systems NF (180 MWCO) and RO</td>
<td>0.0081</td>
<td>–</td>
<td>98.7 vs. 99.7, NF vs. RO</td>
<td>Yes</td>
<td>20 vs. 80%, NF vs. RO</td>
</tr>
</tbody>
</table>
retain the dissolved chemicals whilst removing colour down to very low levels. Ultrafiltration membrane processes allow selective recovery of sparingly soluble dyes, such as indigo, but downstream reverse osmosis is then usually required to remove soluble dyes (Cheryan, 1998; Woerner et al., 1996; Marcucci et al., 2001; Sójka-Ledakowicza et al., 1998). Having said this, some interesting results have been reported by Porter and his various co-workers whereby low molecular weight anionic dyes were found to be substantially rejected by a titanium dioxide microfiltration membrane (Zhang, 1996; Porter and Gomes, 2000). This was attributed to charge rejection, and charged UF membranes have been successfully employed in South Africa for selectively recovering water and dissolved salts (Erswell et al., 1988).

Most operating membrane-based dyewaste recovery and reuse systems ultimately make use of reverse osmosis or nanofiltration to decolourise. Payback periods are invariably long even where freshwater costs are high, although two bench-scale studies have reported calculated operating costs of between $0.5 and $1 per m$^3$ product (Voigt et al., 2001; Ciardelli et al., 2001). For two South Africa installations the payback ranges from just less than three years for an NF process (Buckley, 1992), to over seven years for a system combining ceramic UF with RO (Short, 1993). A payback period of 1.37 years was quoted for a UF/NF plant in North Carolina (NCDENR, 1995), although no further detail was given. Given the continuing downward pressure on membrane costs, the increasing stringency of environmental regulations worldwide and the history of the employment of membranes for textile wastewater treatment and recycling in certain regions of the world, and South Africa in particular, it seems inevitable that the use of membranes for this duty will become more widespread in years to come.

References


Industrial waters 155


Colour Index, 3rd Revision (1987). Society of Dyers and Colourists (UK) and the American Association of Textile Chemists and Colourists (USA).


3.4 The beverage industry

The food and beverage industries are major consumers of water, with the beverage industry in particular consuming as much as 10–12 tonnes of water per tonne of product – or even more for brewing. The majority of water consumed in this industry is used in washing and cleaning operations, and as such represents significant opportunities for reclaim and recycling. Global water usage within the two sectors is difficult to define, but some available data for beverage production (Table 3.35) suggest global usage of around two billion (i.e. thousand million) tonnes of water per annum. The food industry in the USA alone consumes 4 billion tonnes of water per annum – 50% more than the second largest user, the pulp and paper industry (Levine and Asano, 2002).

The food and beverage industries do not generally reuse or recycle water which is either used in or comes into contact with the product. This is primarily a marketing and public perception issue in the same way as recycling of sewage effluent for potable water has significant consumer acceptance problems (Section 1.1). Since a significant amount of water in the industry does not go into the product, opportunities still exist for water reuse and the quality of water required for use in the product is not normally of concern for recycling. However, the water quality demanded for washing the product or product containers purposes is usually of potable standard, and there is still a reluctance to use recycled water even for these duties. Recycled water must be either recycled at the point of use to avoid additional contamination or recycled to non-product uses, such as utilities (usually power generation and heat transfer) and washing. Fortunately, because food, dairy and, in particular, brewing processes are energy intensive, the utility water consumption in boilers and coolers is quite high and can demand up to twice as much water as the primary production process.

3.4.1 Point of use recycling opportunities

Bottle washers

In the dairy and beverage industries the bottle washer is a significant user of water, and most washers have a make-up water flow rate to the final rinse of about 10–20 m³ h⁻¹ (which equates to 250 ml per bottle). A proportion of this water is used to pre-rinse the bottles but a significant part of this final rinse water is discharged to drain. The water is generally low in turbidity with a pH of about 10.5 and a conductivity of about 2500 μS cm⁻¹.

Table 3.35  Global water use. beverage industry

<table>
<thead>
<tr>
<th></th>
<th>Volume, m³ p.a</th>
<th>Global water use, m³ p.a.</th>
<th>Average weight ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soft drinks</td>
<td>380 000 000</td>
<td>1 140 000 000</td>
<td>3</td>
</tr>
<tr>
<td>Breweries</td>
<td>136 000 000</td>
<td>952 000 000</td>
<td>7</td>
</tr>
</tbody>
</table>
Several systems have been used to recycle this water, with various degrees of water product purity. Filtration, pH adjustment and chlorination does not address the high TDS and demands a constant bleed: the treated water quality does not meet potable guidelines and consequently this system has only been used in the event of severe water shortages. Carbon dioxide addition, dealkalisation by ion exchange and UV sterilisation represents an improvement over filtration-based methods and has been implemented in some German facilities. Filtration followed by pH correction and reverse osmosis has been used only in very few installations because of the high capital and operating cost, and can only be justified where both the water and effluent costs are high.

In the majority of cases the water from the final rinse can be used virtually untreated in applications such as crate washers and as pasteuriser make-up after cooling. In addition the water can be collected and used for floor washing. Because the cost of treatment in these cases is negligible, most of the reclaim applications adopt these procedures.

A recent example of water recycling and reuse is at the Coca Cola Amatil plant in New South Wales (Environment Australia, 2001). Two simple recycling initiatives have been undertaken at this plant, the first involving reuse of the backwash water and the second the reuse of container rinse water. Interestingly, the first of these, which recovers around 200 m$^3$ day$^{-1}$ of backwash water from the sand and carbon filters, appears to be blended with the mains water and reused in the manufacturing process. The blend is maintained at less than 1:5 recovered:mains water. The payback time for the recycling system, which comprises pipework and a backwash water recovery tank, is estimated to be around two years. The recovered container rinse water, on the other hand, is used in the evaporative cooling towers following filtration. About 16 m$^3$ day$^{-1}$ is recovered for this duty.

**Caustic recovery**

In the majority of food and beverage applications a large amount of caustic soda is used for bottle washing and CIP (clean in place) applications. The disposal of the spent caustic solution is problematic and expensive. In most cases the effectiveness of the caustic solution is assessed by assaying for carbonate contamination or dirt content; when these levels reach a certain limit the solution is disposed of. The caustic content may still be quite high, however, and nanofiltration membranes have been developed (Koch and PCI Memtech) to clean and concentrate the spent caustic solution. The process plant for this duty is quite expensive, and the economics are such that the plant is only justified if the caustic volume used and cost of its disposal are both very high.

**Bottle or can pasteurisers**

An improperly balanced pasteuriser can use a large amounts of water which is often discharged direct to drain. In most cases this water can be recycled back to the pasteuriser directly after cooling and filtration. Checks must be made on product contamination in the case of bottle pasteurisers.
**Maltings steep water**

A significant amount of water is used as steep water in maltings, where the water is used to soak the barley. The effluent contains a high concentration of organics, making it expensive to dispose of. Trials at several maltings have shown that nanofiltration membranes or some types of reverse osmosis membranes can produce water suitable for reuse (PCI Memtech). The cost of the effluent disposal is not reduced, however, as the organic loading rate from the retentate stream to drain remains the same.

**Milk processing/condensate**

Membranes have been used for many years in the dairy industry for process separation applications (Cheryan, 1998). Across the whole food industrial sector dairy applications probably account for the largest proportion of installed membrane capacity. Indeed, it is the selectivity of the membrane filtration processes, in terms of retentate molecular size, which allows fractionation of milk to produce cream and skimmed milk by microfiltration and protein from lactose by ultrafiltration. Since a key membrane property is its thermal stability, generally to around 50–55°C to permit operation at lower fluid viscosities, and chemical stability, to permit more aggressive chemical cleaning and sanitation, polysulphone, polyethersulphone or PVDF membranes (Table 2.3) are generally used.

The use of membranes for other applications such as condensate recovery is now an established technology. In powdered milk production facilities a large amount of steam is used for evaporation purposes, and the condensate recovered from the evaporators is both hot and relatively pure. This makes it ideal for make-up water for the boilers. To remove the organic contaminants special high-temperature reverse osmosis membranes (Duratherm Excel by Desal) are used and the reverse osmosis plants are equipped with sophisticated CIP systems to clean the units on a daily basis.

**Other applications**

Other membrane applications have been used in various industries for effluent reduction and product recovery. These include cross-flow microfiltration for beer recovery from tank bottoms (Vivendi Memcor) and vibrating membranes for beer recovery from spent yeast processing (Pall VMF®, Section 2.1.4). These applications can normally be economically justified from savings based not on water but on product recovery and reuse.

**3.4.2 End of pipe recovery opportunities**

As mentioned in previous sections, once the effluent has been mixed and the risk of contamination has increased, the potential uses of the water are reduced. Effluent from most UK food and beverage industries is discharged direct to sewer and the associated costs paid. To recycle effluent would first involve an effluent treatment plant, generally employing primary biotreatment for high organic loadings with downstream polishing using depth and/or membrane filters,
possibly preceded by coagulation. In many cases such an option is only marginally economically viable given the current level of effluent charges, cost of plant maintenance and the incentives provided by water companies to keep discharging to drain. However, having treated the effluent to a reasonable level, the costs of treating to a standard suitable for recycling and thereby avoiding water costs often shifts the economics in favour of the plant recycling option.

The economic case is further enhanced if (as in the case of the food and drinks industry) the water is reused in high-quality applications such as boiler feed. This is because the quality of water produced by an RO plant is generally of a higher quality, with reference to key parameters such as hardness, silica and total dissolved solids, than mains water. It is almost certain that the company will already have to treat the mains water separately for boiler use. Often the quality of the RO recycled water will be better than the current feedwater and so that savings in boiler chemicals and heat can be made. When all of these factors are evaluated, the economic case for effluent treatment and recycling may be justified. Several effluent recovery and reuse plant have already been installed in the UK, and it is likely that once the treatment and pretreatment regimes have been established and proved many more will follow.

**General aspects of plant design and operation**

Many plants have used biological treatment plants with filtration and chlorination prior to cellulose acetate-based reverse osmosis. This can work successfully providing the level of filtration (sometimes dual media filtration) is sufficient. The cellulose acetate (CA) membranes are less prone to fouling and can tolerate a chlorine residual (Table 2.3), so biological fouling is reduced. Unfortunately CA membranes are not as widely used within the industry because of higher power costs, and lower rejections and pH tolerance. Composite polyamide membranes are more commonly used and can be used successfully, but pretreatment becomes more critical since they are more prone to fouling than CA membranes. In a few plants employing tertiary media filtration severe problems have been encountered in maintaining the flux within the RO plant. In most cases either a membrane biological treatment process (i.e. a membrane bioreactor, MBR) is required or alternatively ultrafiltration must be used as the basic pretreatment step. The UF configuration and cleaning regime will depend on the upstream process, as determined by pilot trials.

With an effective UF plant or an MBR as pretreatment, the use of polyamide RO membranes should not cause a problem. It is advisable however to use much higher fouling allowances (such as 30–50%), such that the pump pressures are significantly higher than design projections based on osmotic pressure alone. It has been found in some plants that following initial organic fouling the membrane flux drops to a sustainable level. This must be allowed for in the plant design, and can normally be determined by pilot trials.

Biological fouling can normally be controlled by biocide addition. This can be carried out periodically on line using a non-oxidising biocide, but can only be employed if, as in most cases, the water is not being used for potable applications. In some cases chloramine formation has been used successfully to protect the
membranes. In this case pH control is critical, since at higher pH levels chloramine dissociation takes place resulting in damage to the membrane.

References

3.5 Pure waters in the pharmaceutical industry

3.5.1 Background
The pharmaceutical industry is a global one with multinational companies carrying out research and development activities, clinical trials and production on different continents and supplying the products to virtually every country of the world. The industry is relatively unaffected by the “bear” and “bull” markets that typically exert a dramatic effect on other high-technology industries, such as electronics. The market for pharmaceutical products continues to grow as new drugs are developed to treat more and more previously untreatable conditions. This development is being sustained or even accelerated by the new biotechnology-based products passing through clinical trials and coming onto the market.

A pharmaceutical company uses water for many different purposes, many of these being unrelated to the pharmaceutical activities of the company. For such general applications for water, such as for boiler feed, heat transfer, toilet flushing, showering, laundering, fire control, etc., recycle and reuse considerations are no different to those within other industrial sectors: they are applications that require a water feed and generate an effluent stream, and may or may not be suitable applications for utilising recovered water based on water quality, quantity and processing costs. However, there is nothing about these applications that is changed by their being carried out within a pharmaceutical organisation.

Water is used extensively in the pharmaceutical industry and it is the most frequently used ingredient of pharmaceutical preparations. There are a wide range of products and intermediate products that require a reliable supply of water during their manufacture, including:
- bulk pharmaceutical compounds/active pharmaceutical intermediates (APIs).
- over-the-counter (OTC) products – cough medicines, paracetamol, contact lens cleaners, etc.,
- prescription medicines (tablets, ointments, creams, liquids),
- "health" foods (vitamin tables, energy drinks),
- "cosmaceuticals" (antiseptic ointments, concealers),
- aerosols/inhalers, and
- injectable drugs.

The importance of water and water quality was increased enormously by the introduction of parenteral (injection and intravenous infusion) therapy. Water may be a raw material, a process intermediate, a product ingredient, or even the product itself. Its wide use in many areas related to the production and control of medicines demands that manufacturers pay very close attention to process water quality. Further, apart from "mechanical" errors such as mistakes in labelling of products, final product recalls for problems which can be related back to the water used in production of the drug accounts for the next largest group of product recalls.

In planning and operating a pharmaceutical water system the following are a few of the areas that must be considered:

- overall requirements for water,
- specifications and purification methods to be adopted,
- installation and validation of water systems,
- routine quality monitoring requirements, and
- capital and operating costs.

### 3.5.2 Water quality standards

The quality of materials used in the manufacture of pharmaceutical products are all defined in pharmacopoeias. The main pharmacopoeias referred to today are the European Pharmacopoeia (EP), the United States Pharmacopoeia (USP 25) and the Japanese Pharmacopoeia (JP). The standards defined in these pharmacopoeias are all enforceable in law and all manufacturers of pharmaceutical products are regularly subjected to inspection by medicines inspectors from countries where they have a licence to sell their products. These inspections are carried out to ensure that the manufacturers are meeting the quality and production requirements defined in their product licences, as defined in the pharmacopoeias and as required for compliance with current good manufacturing practice (cGMP). A company failing to meet these standards may be given a warning letter from the inspection authorities and a defined length of time to rectify the problems. If this does not take place within the stipulated time limit the company may have their licence(s) withdrawn until such time that the problem is resolved.
Water is unique within all the pharmacopoeias in that it is the only material commonly used by pharmaceutical companies which not only has quality standards defined but production methods are also defined. Two main water qualities are defined as Purified Water (PW, Table 3.36) and Water for Injections (WFI, Table 3.37). The definitions of the quality standards, the production methods for these and the monitoring methods vary slightly between the different pharmacopoeias.

According to the European Pharmacopoeia 2000, Purified Water is water for the preparation of medicines other than those that are required to be both sterile and apyrogenic, unless otherwise justified and authorised. Definitions are given in the EP for PW in bulk and in containers. Purified water in bulk is used as an excipient in the preparation of non-sterile products and as a starting material in the preparation of water for injection and pharmaceutical-grade pure steam. It is also used for rinsing purposes (cleaning of containers) and in the preparation of cleaning solutions. Purified water in containers is purified water in bulk that has been filled and stored in conditions designed to assure the required microbiological quality. It must be free from any added substances. Although very specific stipulations concerning PW quality (Table 3.36) and quality control are given, all process technologies are permitted for its production. In the

<table>
<thead>
<tr>
<th>Table 3.36 Purified Water (PW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>TOC</td>
</tr>
<tr>
<td>Conductivity</td>
</tr>
<tr>
<td>Conductivity</td>
</tr>
<tr>
<td>Nitrate (NO₃)</td>
</tr>
<tr>
<td>Heavy metals</td>
</tr>
<tr>
<td>Aerobic bacteria</td>
</tr>
</tbody>
</table>

* Determined by membrane filtration, using agar medium B.

<table>
<thead>
<tr>
<th>Table 3.37 Water for Injection (WFI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>TOC</td>
</tr>
<tr>
<td>Conductivity</td>
</tr>
<tr>
<td>Conductivity</td>
</tr>
<tr>
<td>Dry residue</td>
</tr>
<tr>
<td>Nitrate (NO₃)</td>
</tr>
<tr>
<td>Heavy metals</td>
</tr>
<tr>
<td>Aerobic bacteria</td>
</tr>
<tr>
<td>Bacterial endotoxins</td>
</tr>
</tbody>
</table>

* Determined by membrane filtration, using agar medium B.
USP 25, there is similarly no stipulation of process for PW production. Organic carbon measurement can be by TOC, a combustion-based instrumental method, or by permanganate value (PV).

Sterilised water for injection is used for dissolving or diluting substances or preparations for parenteral administration. Water for injection in bulk is used in the manufacture of parenteral and ophthalmic products. It is also used for final rinsing of containers (e.g. primary packaging materials) and manufacture of these products. In addition to stipulations concerning quality (Table 3.37) and quality control, the EP 2000 stipulates that the water must be produced by distillation. The USP 25, on the other hand, permits both distillation and reverse osmosis.

Potable water, or water intended for human consumption, is also used as feedwater for the production of purified water. Potable water may be used to rinse product-contacting surfaces of equipment. Treated potable water has the same uses as potable water and water intended for human consumption but has been treated to reduce its microbial content. Finally, highly purified water (non-compendial water) is used in the preparation of medicinal products where bacterial endotoxins need to be controlled, except where water for injection is required. Current methods for the preparation include double-pass reverse osmosis, reverse osmosis combined with ultrafiltration and distillation.

While the purity levels specified for PW and WFI are not as high as those required in industries such as electronics and power station water systems (Table 1.1), the control of the quality and the documentation related to the system are of paramount importance. In an extreme case, if the quality of water is not achieved a patient may ultimately die when being treated with medication manufactured using that water. Consequently, the pharmaceutical industry invests enormous sums of money to "validate" water systems, a process that continues throughout the life of each water system.

Give that control of product water quality is crucial in this industry, control of feedwater quality is similarly important. The USP includes a section that gives general information on water for pharmaceutical purposes. This section describes different types of water used, i.e. Drinking Water, Purified Water, Sterile Purified Water, Water for Injection, Sterile Water for Injection, Bacteriostatic Water for Injection, Sterile Water for Irrigation and Sterile Water for Inhalation. It begins with the statement that the feedwater used for pharmaceutical preparations needs to be of potable water quality, meeting the requirements of the National Primary Drinking Water Regulations (NPDWR) (40 CFR 141) issued by the Environmental Protection Agency (EPA), since this "ensures the absence of coliforms". It is also pointed out, however, that meeting the National Drinking Water Regulations does not rule out the presence of other microorganisms, which, while not considered a major public health concern could, if present, constitute a hazard or be considered undesirable in a drug substance or formulated product.

The stipulation of compliance with the national potable water standards is critical, and effectively severely constrains recycling opportunities. This ensures that the level of impurities, inorganic, organic and bacterial, that can be present
in the water are defined and controlled. In most pharmaceutical facilities municipally supplied drinking water is used as the raw material. In a few facilities local borehole water, or sometimes river water, is used as the starting material but this must be monitored and tested to show it meets the standards required for drinking water.

3.5.3 Volumes and quality of aqueous process and waste streams

Within a pharmaceutical facility there are a wide range of process-related aqueous effluent streams. Firstly there are effluent streams from the water purification process itself. Since these mostly contain the concentrated impurities present in the drinking water supply it is unlikely that such streams would be suitable for recycling within the water treatment system since this would lead to accumulation of impurities, but they may be employed in other site applications. The typical waste streams are back-washings from regeneration cycles on multimedia filters, organic scavengers and water softeners. In a pharmaceutical water system it is common practice to remove hardness from the water by passing the feed water through a cation resin exchanger in the sodium form to exchange hardness ions (calcium and magnesium) with sodium. Consequently, the RO reject water will normally be softened filtered water with a conductivity up to four times that of the potable mains feed. Depending on the feedwater conductivity this water may be suitable for use for similar applications as those of recovered grey water, although it cannot be reclaimed within the water treatment system.

Water that may be suitable for recycling within the water treatment system is that which is sent to drain from a final polishing unit, for example the drain stream from an electrodeionisation (EDI) unit (Section 2.1.4), where the level of concentrated impurities is still less than that present in the mains feed supply. The other source may be from water that is sent to drain when the system is in internal recycle. Unlike many applications, a pharmaceutical system is typically designed to have all water in continuous motion with internal recycle loops on the pre-treatment and the purification sections of the plant as well as the final distribution loop. This is because the biofilm growth varies inversely with water velocity: continuous motion of the water suppresses the creation of bacterial colonies or biofilms on the pipe making it easier to maintain control of the microbiological levels. This high-purity water is recirculated and becomes the feed stream to the RO system, such that the reject water is still of a high quality and would be suitable for reclaim back to the start of the water system. The only potential problem with reclaiming this water is that the heat input from the RO pump would also be recovered, such that the temperature of water in the system would gradually increase. This may ultimately result in either a cooling unit or dumping of water to reduce the temperature.

Clean in place (CIP) cycle effluent, depending on the type and stage of the CIP process, may contain significant quantities of product residues and detergents from the first rinse cycle or else could contain virtually pure water from final rinse. Most CIP systems employ a series of cleaning procedures operated at
different temperatures and/or using different cleaning chemicals depending on
the residues that need to be removed from the equipment. All cleaning cycles
normal finish with one or more rinses using drinking water then a final rinse
using PW or WFI.

Finally, water may be “dumped” from the system periodically as part of a
standard procedure. Until quite recently it was standard practice for all WFI to be
“dumped” after 24 hours if it was not used in production so that all water could
be classed as “freshly produced”. This procedure is not commonly followed now
but where it is carried out the water being dumped is of a much higher quality
than the drinking water used to produce more WFI. Water is also dumped to
regulate the load on the sanitiser where sanitisation by heating to 80°C is
conducted. Such water is also of a much higher quality than the mains water
used as the feed for the purification system.

In the UK and Ireland the majority of the PW and WFI water systems produce
water at between 500 and 5000 l h⁻¹, although there are some notable
exceptions where generation rates are significantly higher. There is often a PW
and sometimes a WFI water system installed in each building on a large site: one
UK pharmaceutical site has approximately 30 PW and WFI water systems on a
single site. On smaller sites one generation plant can supply a number of
buildings or manufacturing areas via distribution pipework systems.

Since every production process is different, requiring different volumes of
water, producing different quantities of effluent with different levels and types
of contaminant, it is not possible to estimate typical water usage figures.
However, to illustrate the water volumes that might typically be involved a
hypothetical system can be considered.

PW and WFI systems in Europe are typically based on generation flow rates of
0.5–5 m³ h⁻¹. There are some notable exceptions utilising higher flow rates and
typical system sizes in America also tend to be higher. This hypothetical system is
based on a PW production rate of 3000 l h⁻¹ and 1000 l h⁻¹ WFI, with a daily
process usage of 24 000 l of PW and 8000 l of WFI. The PW system will operate
continuous in recirculating mode when no water is being made up to the storage
vessel while the WFI still will operate in a stop/start mode. The approximate
quantities and qualities of effluent will be in the ranges shown in Table 3.38.

Based on these approximate estimated volumes the maximum volume of
effluent water that could be reclaimed is around 31 m³ day⁻¹. If the water were
passed to a purpose-built reclaim system this volume would rise to around 70
m³ day⁻¹. In practice the water that arises from some of the more contaminated
CIP rinses would probably not be reclaimed.

If the main technology used in the reclaim system is reverse osmosis then
the likely maximum quantity of reclaimed water which could be returned to the
purification system as “raw” water would be around 23 m³, based on a recovery
of 75% from the RO treatment plant. Based on current costs of around £1.6 per
m³ ($2.57 per m³) for purchase of potable water and disposal of effluent water,
reclaiming this quantity of water would save only around £37 [$59.5] per day.
The operating costs associated with the reclaim system, including maintaining
the validation documentation necessary, is estimated to be over £50 [$80.4] per
Table 3.38 Representative data for effluent water generation

<table>
<thead>
<tr>
<th>Source</th>
<th>Flow (m³ day⁻¹)</th>
<th>Comments/assumptions</th>
<th>Effluent water quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>PW generation</td>
<td></td>
<td>RO reject while water is made up to PW storage vessel</td>
<td>1200 µS cm⁻¹</td>
</tr>
<tr>
<td></td>
<td>11</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>13</td>
<td>RO reject while system is on recycle</td>
<td>75 µS cm⁻¹</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>EDI waste stream</td>
<td>200 µS cm⁻¹</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>Pretreatment regeneration/backwash</td>
<td>High salt and contaminant content</td>
</tr>
<tr>
<td>Total</td>
<td>31</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CIP, PW</td>
<td>12</td>
<td>Assume half of PW is used for final clean rinse stage</td>
<td>5 µS cm⁻¹</td>
</tr>
<tr>
<td>CIP, process water</td>
<td>24</td>
<td>Initial cleaning cycles – main contain residual product, cleaning chemicals, etc.</td>
<td>300–1500 µS cm⁻¹</td>
</tr>
<tr>
<td>Total</td>
<td>36</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WFI</td>
<td>0.8</td>
<td>10% blowdown over 8 hours</td>
<td>10 µS cm⁻¹ at 90°C</td>
</tr>
<tr>
<td>CIP, WFI</td>
<td>2</td>
<td>From the final rinse stages using WFI water</td>
<td>5 µS cm⁻¹</td>
</tr>
<tr>
<td>Total</td>
<td>2</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Feed water conductivity assumed to be 300 µS cm⁻¹.

*a* Candidate water for reclaim within water treatment plant: 30.8 m³ day⁻¹ in total, 54 µS cm⁻¹.

day based on electrical usage, maintenance labour, microbiological testing, chemical testing and validation activities. The operating cost of the reclaim system would thus almost certainly exceed the potential saving, even without considering recovery of capital cost, for a small system such as this. Consequently, even without the barrier imposed by the pharmacopoeia specifications and the views of government inspectors prohibiting the use of any starting material other than drinking water, there is unlikely to be an economic justification for reusing effluent water from the process. Any such water would need to be combined with other effluent water from non-process applications on site to make a dedicated reclaim system economically attractive. On the other hand, reuse of some of the more dilute streams, specifically some of the CIP streams and any other high-purity effluent streams within the existing purification scheme would be much more likely to be cost effective.

### 3.5.4 Reuse opportunities

There appear to be no examples of any facility using recycled plant/process effluent water treated to bring it up to the quality of drinking water and then being used as the feed stream to produce PW or WFI and, under the current regulations, there are no obvious reuse opportunities in the pharmaceutical sector. One UK facility, based in a very remote rural location, submitted a proposal to the Medicines Control Agency (MCA) in the UK for a facility that included effluent recycling plant and they were told that this was not acceptable to the inspection authorities. Since no local effluent facilities existed all their plant effluent had to be collected and removed from site by tanker because of this decision. This situation, which is global, may change if regulations are relaxed.
Membranes for Industrial Wastewater Recovery and Re-use

Table 3.39  Highly purified water (non-compendial water)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>USP 25</th>
<th>Ph. Eur. (bulk)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOC</td>
<td>ppb C</td>
<td>-</td>
<td>500</td>
</tr>
<tr>
<td>Conductivity</td>
<td>µS/cm at 20°C</td>
<td>-</td>
<td>&lt;1.1</td>
</tr>
<tr>
<td>Aerobic bacteria</td>
<td>CFU/ml</td>
<td>-</td>
<td>&lt;10</td>
</tr>
<tr>
<td>Bacterial endotoxins</td>
<td>EL/ml</td>
<td>-</td>
<td>&lt;0.25</td>
</tr>
</tbody>
</table>

and costs of water and effluent disposal increase, the latter being almost inevitable.

It is nonetheless technically possible to reuse effluent at present to generate a usable feed stream, especially given that around 45% of the wastewater is likely to have an aggregate conductivity of well below 100 µS cm⁻¹, based on the data given in Table 3.39, and would thus actually dilute the mains water supply. Much of the effluent produced from pharmaceutical processes has a limited range of contaminants, namely only trace levels of the products being manufactured and very low levels of salts and organics. Also, this effluent is much better characterised and controlled than typical municipal-supplied drinking water, since PW and WFI waters are very low in dissolved matter and the exact nature of the impurities added are known. Consequently, it should be possible to specifically monitor and, if necessary, selectively remove individual impurities introduced from the production environment into the effluent to ensure that the reclaimed water consistently meets the drinking water standard.

For example, PW typically has a conductivity < 1 µS cm⁻¹, i.e. a total dissolved solids concentration of around 0.5 mg l⁻¹. This content is largely made up of sodium and chloride ions with traces of potassium, fluoride and other small molecular weight ions. If this water is used in the final rinse of a process vessel the only contaminants added will be residual quantities of the products being manufactured, generally at levels < 1 mg l⁻¹. The actual quantities of these products will be accurately measured and recorded as part of the validation of the CIP process but will obviously vary between applications. The impurities from other waste streams will generally be the same as those present in the feed water but at much lower levels, and can be measured using the same analytical methods as are used for potable water.

Water continues to be the most commonly used chemical in most pharmaceutical operations and also continues to be one of the most problematic. One of the largest causes for product recalls are water-related issues, and such an environment imposes a considerable barrier to water recycling. If there is also no strong economic driving force, regardless of the technical feasibility of water recycling, it is unlikely to happen in this industry in the near future.
Chapter 4

System design aids

4.1 Computer-aided design for reverse osmosis plant
Etienne Brauns
VITO

4.2 Water pinch analysis
Danielle Baetens
VITO

4.3 Design examples
Simon Judd
School of Water Sciences, Cranfield University
4.1 Computer-aided design for reverse osmosis plant

4.1.1 Introduction

It has already been pointed out (Section 2.3.2) that modelling of filtration processes is not feasible on a universal basis, and that only reverse osmosis can be modelled on the basis of solution–diffusion. In such a case, modelling is possible using simple analytical expressions provided the water can be treated as a continuum essentially free of the more complex interactions introduced by the presence of dispersed particles, solute precipitation effects (gel layer formation) and/or biological activity at the membrane–solution interface. As a result of these complicating factors, which relate largely to porous membrane processes, computer-aided design (CAD) software is currently available only for reverse osmosis membranes and modules. The software packages are based on a series of calculations that enable the estimation of the RO array design and operation. It is instructive to review these CAD packages, which are produced by RO membrane manufacturers and suppliers based on their own products, with a view to ascertaining their general usefulness and limitations (Brauns, 2001).

4.1.2 Key elements of the reverse osmosis process

Modelling of the reverse osmosis process relies on accurate mathematical representation of:

- the osmotic pressure,
- concentration polarisation, and
- the hydraulic resistance offered by the membrane and the membrane channels.

As already discussed (Section 2.3.1), the pressure required to drive a reverse osmosis process relates both to the permeability of the membrane and the osmotic pressure. The osmotic pressure is solute concentration related, and can be very high for highly saline solutions: the osmotic pressure for seawater (35 g l\(^{-1}\) NaCl) is around 27 bar. No flow of water takes place unless this pressure is exceeded. At moderate concentration values, osmotic pressure varies almost linearly with concentration for many univalent salts, including sodium chloride, according to the van’t Hoff equation (Chapter 2, Equation (2.9)). In this equation \(y\) depends on the degree of dissociation of the salt. Non-linear correlations of \(\Pi\) with \(C\) have been fitted to Taylor series and power relationships:

\[
\Pi = RT(y_1 C_1 + y_2 C_2 + \ldots) \quad (4.1)
\]

\[
\Pi = yRTC^n \quad (4.2)
\]

Any one of the above relationships may be employed in existing RO design software.
Concentration polarisation (CP) can be quantified from film theory (Section 2.3.2, Equations (2.14)-(2.18)), and its effect on osmotic pressure and scale formation respectively determined from the van't Hoff equation (or the above modifications thereof) and equilibrium thermodynamics expressions for precipitation (Section 2.4.3). Whilst the effect on osmotic pressure can be calculated with reasonable accuracy, the actual scale deposition rate cannot. This is because the simple expressions outlined in Section 2.4.3 are derived from equilibrium thermodynamics and cannot account for the very wide variations in the scaling kinetics that exist. Indeed, it is quite common for some scalants to remain dissolved at concentrations well beyond the theoretical saturation limit determined from the solubility constant $K_s$ (Table 2.14), in a supersaturated state, even without the addition of threshold inhibitors (Table 2.15) to suppress nucleation.

In principle, film theory also permits the optimisation of both the retentate flow rate and the design of the spacer material in the spiral-wound RO element. However, in practice the choice of RO spacer material is dictated to a large extent by cost and, in any case, is not one that can be made by the RO process designer but by the membrane supplier. Similarly, although turbulence, and hence mass transfer, is promoted by a high cross-flow velocity, a practical limit is placed on the retentate flow rate by the construction of the spiral-wound element, which can only withstand a certain maximum hydraulic loading without risk of failure. Failure can be manifested through a phenomenon known as “telescoping”, where the centre of the element is pushed axially outwards by the force of the flow. Since the specific resistance (i.e. the hydraulic resistance per unit length) offered by the membrane channel is constant along the length of the membrane element, the limit on retentate flow rate implies a limit on both the inlet pressure and the pressure drop across the element. For large elements, 1 m long, the maximum pressure drop is normally much less than 1 bar.

Most RO filtration systems are based on the spiral-wound configuration, a noteworthy exception being the Monsanto Prism hollow fibre membrane. The largest of the spiral-wound elements are 8 inch (about 0.2 m) in outer diameter and 40 inch (about 1 m) in length, with newer membrane elements being 1.5 m long. Membrane elements are mounted inside pressure vessels in which they are connected sequentially to give a module length of up to 8 m or more for an 8-element module. These modules can then be arranged in parallel and/or in series to produce a matrix, or array (Section 2.4.1), of membranes with a sufficient total membrane surface area to obtain a specified total permeate flow and permeate quality at an acceptable cost.

Both the appropriate number of elements in the module and the arrangement of modules in the array to obtain volumetric flow information can be arrived at through a simple mass balance (Section 4.3.2) provided the conversion $\Theta$ per membrane element can be assumed constant. However, since (a) $\Theta$ must necessarily be a function of both the transmembrane pressure (TMP) and the osmotic pressure $\pi$, and so of the solute concentration, and (b) the solute flux through the membrane and the fouling propensity are both critically important, a simple water balance is normally insufficient. To obtain water quality
information a mass balance of the solutes is necessary. This is considerably more complex than the water balance solution, which is obtained analytically using Equation (2.22), because of the interrelationship between TMP, Θ and π, and the complicating influence of concentration polarisation (Section 2.2.3). This means that the impacts of the individual design parameters on operation (Table 4.1) are largely interdependent, and that the array design solution is then only obtainable iteratively.

To enable these array design calculations to be carried out, RO design software packages are available for specific RO membrane products. Such software enables the designer to simulate operation of any RO array under any set of operating conditions. These packages (Table 4.2) are intended to allow the user to produce an RO array design from base information, such as the required product flow rate, overall process conversion and feedwater characteristics. The membrane suppliers offer their own RO software to the (prospective) customer gratis. Specific software from universities is also available, or else is under development, at institutions such as RWTH Aachen and Saarland University in Germany, University of Wales Swansea in the UK and the European Institute of Membranes at the University of Montpellier in France. Some of these groups are developing software for nanofiltration.

The starting point for all design packages is the feedwater composition. Whilst data for the principal ions may be available, this is not always the case for some scalant components, such as barium or silicate. These ions form extremely insoluble precipitates, such that they may be close to saturation even at very low concentrations. Most feedwaters, and industrial effluents in particular, are subject to temporal fluctuations in composition, and this has to be accounted for

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Impact</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feedwater composition</td>
<td>Osmotic pressure, scaling, membrane stability, permeate quality</td>
</tr>
<tr>
<td>Feedwater pH</td>
<td>Scaling, membrane stability</td>
</tr>
<tr>
<td>Feedwater temperature</td>
<td>Flux, membrane stability</td>
</tr>
<tr>
<td>Permeate flux</td>
<td>Transmembrane pressure, concentration polarisation, fouling, pressure loss</td>
</tr>
<tr>
<td>Cross-flow velocity</td>
<td>Concentration polarisation, fouling, pressure loss</td>
</tr>
<tr>
<td>Membrane material</td>
<td>Membrane stability, permeate quality, fouling</td>
</tr>
</tbody>
</table>

a Primary or most important in bold type.

<table>
<thead>
<tr>
<th>Company</th>
<th>Membrane name</th>
<th>RO software</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dow</td>
<td>Filmtec</td>
<td>Rosa</td>
</tr>
<tr>
<td>Hydranautics</td>
<td>Hydranautics</td>
<td>Rodesign, Rodata</td>
</tr>
<tr>
<td>Koch</td>
<td>Koch (Fluid Systems)</td>
<td>Ropro, Costpro</td>
</tr>
<tr>
<td>Osmonics</td>
<td>Osmonics, Desal</td>
<td>WinFlows</td>
</tr>
<tr>
<td>Toray, Ropur, Trisep</td>
<td>Toray, Ropur, Trisep</td>
<td>WinCarol, 2p flows</td>
</tr>
</tbody>
</table>
when entering data into the design package. It is also invariably the case that experimentally-determined ionic composition data do not yield an electrically balanced specification. RO design software packages tolerate, to a certain extent, an ion imbalance in the feed composition. Exceeding some predetermined limit of this imbalance normally prompts a request to artificially adapt the total ion balance by compensation (anion or cation). The total ion balance is then usually corrected through the addition of sodium cations or chloride anions, since these ions do not have a substantial impact upon the RO process design and operation. Aggregate aspecific determinants, such as total organic carbon and total dissolved solids, cannot normally be usefully entered into the design package. Most design packages allow entry of data for the most common scalants, such as the sulphates or carbonates of calcium, magnesium, barium, strontium and iron, but not the more unusual compounds, such as sulphides of many divalent metals, that can arise in some industrial effluents (Table 2.14).

The composition of the feed determines both osmotic pressure and scaling propensity. These are increased both by the conversion, the ratio of permeate to feed flow, and the degree of concentration polarisation (CP). In most software packages a limit is placed on the CP coefficient (normally denoted as $\beta$-coefficient). For example, for the RoPro package produced by Koch-Fluid Systems the maximum $\beta$-coefficient value is 1.1. This limit is based on experience, rather than anything derived from first principles. A warning is normally given when the retentate scalant concentration exceeds its equilibrium solubility limit, demanding remedial measures in the form of appropriate pretreatment (Section 2.4.3) or else reduced flux.

Other specific feedwater quality determinants of importance are the pH and temperature. The feed pH is of particular significance since it has an impact upon both scaling propensity and membrane integrity (see below). Precipitation of hydrolysable scalants is suppressed at low pH levels and most RO CAD packages allow for pretreatment with acid or base. The feedwater temperature has a direct impact upon the permeate flux through the viscosity (Section 2.3.1), as well as on scalant solubility and biological growth. Calcium carbonate solubility and biological growth both decrease with increasing temperature, whereas the viscosity (and hence the overall permeability) increases. Since the latter is normally the most important temperature effect, it is crucial, when designing the plant, to base the design on the lowest temperatures likely to be encountered.

Membrane stability has already been discussed (Section 2.1.3). Reverse osmosis membrane materials are all polymeric and have varying levels of chemical stability. The two membranes materials most frequently employed are cellulose acetate (low cost, chlorine resistant but limited rejection and subject to alkaline hydrolysis at pH levels above 6) and aromatic polyamide (high rejection, resistant to hydrolysis but more expensive and susceptible to oxidation by chlorination). The choice of both the membrane material and module will obviously depend upon the application, since parameters such as requirement for sanitisation, feedwater temperature and pH fluctuations and fouling propensity can be critical.
The system hydraulic resistance has a significant impact on the specific energy demand (i.e. the energy consumption per unit permeate product volume). High retentate flow rates reduce concentration polariation, and hence may be expected to increase the flux, but also increase the pressure drop along the length of the membrane element. A high retentate flow rate is associated with a large transmembrane pressure (TMP), and hence a higher flux and recovery. However, this results in an increase in CP. Clearly, there is a balance to be struck between the pressure and the membrane area requirement, such that the flux is not unreasonably high at the module inlet and the cross-flow velocity not reasonably low at the outlet.

It is normal practice to provide a high membrane area by linking several membrane elements in a single module, and by staging (Section 2.4.1) when the retentate flow decreases by 33 or, more usually, 50%. Many RO plant comprise so-called “2-1” arrays, where the flow entering the second stage of the array is half that of the first stage. Staging reduces the membrane area requirement but can also lead to substantial retentate pressure losses. An intermediate pump may then be required to increase the feed pressure to the second stage: this process is sometimes referred to as a “two stage concentrator” where, in this instance, the stage may include sub-stages.

Another way of increasing the overall recovery is to recycle a fraction of the retentate to the feed inlet port of the bank, producing what is normally referred to as a “single pass with recycle” process configuration. This increases the retentate flow rate as a result of combining the recycle stream with the feed stream, such that it is possible to increase the recovery while still maintaining the same degree of concentration polariation. However, the increase in the retentate concentration has a negative impact on both the scaling propensity and the permeate water quality. It is also possible to treat the permeate from the first stage, producing a “twin pass” system (Fig. 2.21).

4.1.3 RO design software

The design of an RO plant involves the adequate specification of parameter values and an appropriate choice of process design. As a result of the multitude of parameters, complex interactions and different design variants, it is not possible to produce RO software able to automatically produce the ideal RO design for a specific application. The design process essentially proceeds by trial-and-error through selection of different types of membranes, design variants and operating parameter values, with the software package indicating when the selected design is unreasonable and (usually) identifying the cause. The packages therefore all demand a rudimentary understanding of the RO process and the effect of the design parameters. Provided this is the case, the packages are normally quite user-friendly and perhaps even didactic in construction. Most have an input “wizard” that guides the user through each of the design elements in sequence. The individual inputs are listed below.
Feedwater quality
The quality of the feedwater(s), comprises composition, pH and temperature, a typical window (from Rodesign by Hydranautics) being shown in Fig. 4.1. Most design packages allow entry of concentrations as either mg/l (i.e. ppm) or meq/l with some also incorporating “ppm as CaCO₃”. Only the most common ions are included, however. Hence, any other ions that need to be added to the feedwater specification must have their permselectivity properties entered. If no such rejection data is available, and they can only realistically be obtained through experimental measurement, then there is nothing to be gained by adding them to the specification. All major suppliers’ packages will prompt if correction of the analysis with sodium or chloride is required to attain electroneutrality. A number of different feedwater streams can be specified for most of the design packages.

Flows and conversion
Flow rates and/or the overall conversion is also a required input (Fig. 4.2, from RoPro by Koch-Fluid Systems). Flow rates can be entered in both imperial and metric units, and it is usual to specify one of the flow rates (feed or product) and the recovery. Some packages allow for bypassing of the membrane array by part of the feedwater. Most of them have a simple percentage fouling allowance that the user can enter to correct for a lower, more conservative estimate of permeate flux when calculating the design.
Process design variant
The process design variant (single pass, twin pass, etc.) normally has to be specified at the start of the design process, and can be selected from a drop-down menu or a dialogue window. The default selected is always the simple single-pass plant. Obviously, the more complex the process design, the more information required to be entered. Fig. 4.3 shows a two-stage concentrator RO plant dialogue window.

Array design
The core of the RO process is the design of the array, in which the number of stages and trains (i.e. rows of modules) must be defined. The definition of a single stage, or bank, includes the number of pressure vessels (i.e. modules), the number of membranes in each vessel and the membrane type. The latter can range from low-selectivity, low-cost softening membranes to the more permselective membranes used for seawater desalination. It is not possible, and not sensible, to specify more than one membrane type in a single pressure vessel or bank, although different membranes may be selected for different stages. Membrane type is selected from drop-down menus that, of course, offer membrane materials specific to the membrane supplier. In some cases, such as ROPRO by Koch, the user can enter an approximate design flux, such that the array can be calculated from this figure, the overall recovery and the properties of the membrane selected. Otherwise, a preliminary value for the number of elements per module can be calculated from the overall required permeate flow.
and the nominal permeate flow per membrane, as specified by a RO CAD package’s specific dialogue window or simply the membrane specification sheet. The element specification can be quite detailed: The Osmonics Winflows package (Fig. 4.4) allows stipulation of element age to account for deteriorating hydraulic performance with age.

**Pretreatment and post-treatment**

Pretreatment selection is usually limited to the selection of acids (sulphuric or hydrochloric) or bases (caustic soda) to adjust the scaling index value (Fig. 4.5), according to the principles outlined in Section 2.4.3. It is normal practice to ignore pretreatment for the first trial run of the CAD and then add the appropriate amount of acid or base according to the scaling index value automatically calculated by the software. Similarly, post-treatment, which can take the form of pH adjustment and/or degassing to remove the dissolved CO₂ down to some selected value, is usually carried out following the first run.

When all necessary input values have been entered, the software is ready to perform a first calculation. The software calculates and presents data for the composition of the different sub-streams plus all hydraulic data (flow and pressure) for all parts of the array (Fig. 4.6). The designer is then able to analyse the results and adjust the operating conditions so as to optimise the system and/or attain the target permeate water quality. This may demand several runs, and the solution ultimately obtained through iteration. The software packages produce a dialogue box indicating warnings (amounting to errors or
incompatibilities in the design), and these can be attended to directly. For example, if the solubility limit of calcium carbonate has been exceeded, acid may be added as pretreatment. If the membrane module is hydraulically overloaded more membranes can be added. If the design selected is entirely inappropriate, calculation of the parameters (which relies on convergence) may not be possible and the software will then produce a warning and halt.

An optimised RO system design is generally considered to be one for which the lowest specific energy demand is incurred and/or the lowest membrane area is demanded at the highest recovery, although these parameters obviously imply a trade-off between operating and capital costs. Although there is no one method of performing the iterative procedure to obtain the optimised system, a few general points can be made:

(i) A high CP level, always greatest at the module exit, should be avoided: a warning is given when the $\beta$-coefficient value in the last membrane of a pressure vessel exceeds the pre-set maximum value. CP can be reduced by increasing the cross-flow velocity and/or reducing the flux. A number of options exist in this regard:

- reduction of overall recovery
- addition of a stage
- increase of feed flow
- reduction of the number of pressure vessels per bank whilst increasing the number of membranes per pressure vessel
- recycling of part of the concentrate
- decreasing the element size.

(ii) Permeate quality changes with recovery. In principle, solute diffusion increases with higher recoveries but permeate flux also increases. These synergistic phenomena mean that permeate quality can increase until some specific recovery value, beyond which it may decrease again. It is rarely the case that the highest permeate water quality, as may be demanded by, for example, microprocessor applications, coincides with the highest recovery.

(iii) The specific energy demand increases with increasing recovery.

(iv) Calcium carbonate scaling can be very substantially reduced through pretreatment by acid dosing or hardness removal, allowing much greater recoveries and thus demanding smaller membrane areas. Hardness reduction can be achieved by classical chemical softening, by ion exchange or lime dosing, or by blending the feedwater with some of the permeate product.

(v) Scaling by non-hydrolysable scalants (Table 2.14) can be suppressed through the use of appropriate scale inhibitors. Dedicated software from anti-scalant chemical suppliers is available (Fig. 4.7) and can be used in parallel with the RO software to calculate scaling propensity of the chemically treated water. Scaling from these contaminants is otherwise reduced only by their removal or by reducing CP.
4.1.4 Cost calculation

Some RO software packages (Koch, Hydranautics) include a provision for costing, which extracts such cost items as capital costs, pumping energy demand, membrane replacement and chemical dosing, automatically from the design file. Pumping energy costs, for example, relate directly to the system hydraulics. Other specific data, such as labour costs, anti-scalant and cleaning chemical unit prices, amortisation, overheads and maintenance, must be entered by the user. Figure 4.8 illustrates the CostPro software (Koch), which permits fairly comprehensive costing of the RO design. Care must be taken when using such packages, since the cost can be sensitive to assumptions made about such things as membrane life, period of capital amortisation and interest rate. On the other hand, the CostPro software allows a direct cost comparison of two candidate designs.

Cost is also very dependent on the scale of operation. Higher specific costs, i.e. cost prices per m³ permeate product, result from smaller plants due to the relatively high capital and maintenance costs, the latter pertaining to labour. A more useful basis for specific cost may thus be the cost per unit time, which is inversely related to the amortisation period and increases with increasing interest rate. Given the sensitivity of costs to maintenance requirements, cleaning frequency, membrane and plant life, interest rates and residual value, it is prudent to compute costs based on a range of assumed values for these parameters which are, of course, largely beyond the designer’s control.
4.1.5 Overview

Whilst RO CAD packages have become increasingly sophisticated and also more user friendly, more versatile and more widely available over the years (Table 4.3), it should be stressed that they cannot provide an alternative to pilot trials. Indeed, all suppliers emphasise that their software is intended to expedite design, rather than provide absolute values for the key process performance parameters with respect to the system hydraulics and permeate product water quality. None of the existing design packages are able to predict organic fouling or equate this, even by approximation, to some measurable feedwater quality determinant such as TOC. It is such determinants that have an impact upon crucial parameters as the pretreatment requirement, frequency and nature of membrane cleaning and membrane life. On the other hand, these CAD packages provide the option of designing a complete RO plant under what could be considered optimum conditions of zero permanent fouling. In practice a more conservative process design and/or operation may well be called for.

One obvious constraint on the use of commercial software packages is their limitation both to reverse osmosis and the suppliers' own products. Several generic RO, and more recently NF, design software packages have been developed over the years by academic and independent research groups worldwide. An example of a pragmatic MS Visual Basic™-based software approach to handle the design and cost calculations for single pass cross-flow, feed and bleed and semi dead-end filtration systems, applicable to all types of...
membranes, is that of Vito. Three (β-version) programs have been written, creating an interactive environment with predefined dialogue windows and allowing the user to handle in a systematic way the numerous filtration parameters (Brauns et al., 2002). The software allows the calculation of both the design of the installation and its global cost or cost per unit permeate product. The absence of accurate universal membrane filtration models, with the obvious exception of RO, is compensated for by enabling the input of basic filtration values, such as, for example, permeate flux or feed pressure, from a datasheet. As a result the software user is able to implement in a suitable spreadsheet a preferred calculation (model) or extrapolation method based on real experimental data to produce basic filtration input values that later can be imported into the Visual Basic® program. This pragmatic approach allows the user to provide the appropriate basic filtration data, and is totally flexible with regards to membrane process and technology. On the other hand, it relies on available hydraulic data (i.e. flux vs. TMP correlations, pressure loss data, etc.) to be usable.

Given the current widespread activity in membrane process modelling, it seems likely that more CAD packages for porous membrane processes will be
## Table 4.3 Summary of RO CAD package capability

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Fluid Systems ROPRO</th>
<th>Film-Tec ROSA</th>
<th>Hydranautics RODesign</th>
<th>Osmontes Winflows</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feedwater and ions</td>
<td>K⁺, Na⁺, NH₄⁺, Ba²⁺, Ca²⁺, Mg²⁺, Sr²⁺, Cl⁻, F⁻, HCO₃⁻, NO₃⁻, SO₄²⁻, SiO₂⁻</td>
<td>Fe²⁺, Mn²⁺, CO₃²⁻</td>
<td>Fe²⁺, Mn²⁺, CO₃²⁻</td>
<td>Fe²⁺, Mn²⁺, CO₃²⁻</td>
</tr>
<tr>
<td>Other constituents</td>
<td>Fe²⁺, Mn²⁺, CO₃²⁻</td>
<td>Y</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Addition of further</td>
<td></td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>constituents?</td>
<td></td>
<td>Blending?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Process design</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Designs</td>
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<tr>
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<td></td>
<td>Two-st.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>concentrator</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Array design</td>
<td></td>
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</tr>
<tr>
<td>Different elements per</td>
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<td>N/Y</td>
<td>Y/Y</td>
<td>N/Y</td>
</tr>
<tr>
<td>bank/stage?</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Design flux input?</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Pretreatment</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acid/base</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Other anti-scalants?</td>
<td></td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Post-treatment</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acid/base</td>
<td></td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Degassing</td>
<td></td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Design limits</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>β_{max} (CP parameter)</td>
<td>1.13</td>
<td>Not given</td>
<td>1.20</td>
<td>Not given</td>
</tr>
<tr>
<td>ΔP_{loss, max (telescoping)}</td>
<td>Membrane specs</td>
<td>Membrane specs</td>
<td>Membrane specs</td>
<td>Membrane specs</td>
</tr>
<tr>
<td>ISI indication/warning?</td>
<td>Membrane specs</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Nominal flux</td>
<td>Membrane specs</td>
<td>Membrane specs</td>
<td>Membrane specs</td>
<td>Membrane specs</td>
</tr>
<tr>
<td>Wizards</td>
<td>Membrane specs</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Input wizard?</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Array wizard?</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Costings?</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Help files</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Data referencing?</td>
<td></td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
</tbody>
</table>

Commercialised within the next few years. As with most CAD packages, these will rely on experimental data for validation purposes and their usefulness is likely to be closely linked to the extent to which they can represent fouling. Whilst immense progress has been made in understanding fouling mechanisms over the last 20 years or so, successful prediction of fouling in porous processes through a consideration of the operating conditions and the membrane and feedwater matrix characteristics is largely limited to model, single-component feedwaters.
4.2 Water pinch analysis

4.2.1 Introduction

As is apparent from the example industrial sectors discussed in Chapter 2, the use of water within and its discharge from an industrial site can be very complex. Minimising water consumption is obviously desirable on environmental and economic grounds, but it becomes very difficult to determine the extent to which this can be done without endangering product quality. The complexity is compounded not only by ignorance of water volumes involved, but also of water quality with respect to key pollutants. The latter can relate both to effluents generated and process water quality demanded by specific processes. Such basic issues have to be addressed ahead of any assessment of water purification technologies, since such purification may not even be necessary. Moreover, the problem of assessment of water demand, discharge and quality is generally compounded by the disparate knowledge base for individual water-consuming processes throughout the company. Clearly an overview is required encompassing all unit operations demanding and discharging water throughout any one site. This information can then be used to form the basis of what is referred to as pinch analysis which, when applied to water systems, is usually referred to as water pinch.

In the following sections the water pinch concept is discussed in detail, and practical methods for its application outlined. Water pinch provides a means of determining the theoretical minimal overall water consumption for a site with different processes, as well as indicating the requirements of a purification technology based on a specific scenario. Although the concepts of water pinch might seem relatively simple, applying them under real process constraints demands a comprehensive knowledge of the overall industrial process to which it is applied. Moreover, as many parameters are involved and interrelationships between the various process streams can be very complex, it is apparent that dedicated software is necessary to expedite the application of the methodology. The latter is widely available although, as with the reverse osmosis software discussed in Section 4.1, the software is merely a tool rather than providing a process solution.
4.2.2 Water pinch: the history

The pinch concept was developed at the Department of Process Integration at the University of Manchester Institute of Science and Technology (UMIST), UK, in 1970 as a method to reduce energy demand by recovering or transferring heat by employing heat exchangers at critical junctures of a process. This pinch concept was then taken and applied to processes using water. The fundamental theoretical formulations for the application of the pinch concept to wastewater problems were amongst others pioneered by El-Halwagi and co-workers (1992, 1995), Smith and co-workers (1991a,b, 1994, 1996) including Kuo and Smith (1997, 1998), Alva-Argáez et al. (1998a,b) and Wang and Smith (1994a,b, 1995). A large number of studies applying the concept of water pinch have since been performed. Mathematical programming approaches have been formulated (e.g. Rossiter and Nath, 1995; Doyle and Smith, 1997), and several software tools are now available (Section 4.2.3). Two such software tools are WaterTarget®️, a commercial software provided by Linnhoff March—a division of KBC Process Technology Limited—and Water, a package provided by UMIST to the members of its own research consortium. In the following sections, these software tools will also be referred to as the LM and UMIST software. WaterTarget®️ is a software suite comprising WaterTracker®️️ and WaterPinch®️️. Whereas the latter is the heart of the program for defining the optimal water network, the first is used to set up the water balance. In the following sections, the program will be referred to as WaterTarget®️️, although WaterPinch®️️ is the part used and elaborated on. The graphical plot used in WaterTarget®️️ does not represent concentration versus mass load, but concentration versus flow rate. This methodology and the interpretation of the curves generated can be found in Buehner and Rossiter (1996).

4.2.3 Methodology

Every problem definition begins with accurately identifying every unit operation using and producing water, including processes as well as utility operations such as steam production. The existing water network is thus obtained and, for given measured flow rates, the water balance can be checked. The accuracy of this balance determines to a great extent the result and usefulness of a pinch analysis. Indeed, a substantial imbalance of water would strongly indicate either unaccounted for water-consuming unit operations, leaks, and/or an ignorance of flow rates through some or all of the selected units. A water pinch analysis can only usefully proceed if the imbalance is less than 10%.

Relaxing the inlet and outlet concentration

It is most often assumed, in the first instance, that all processes are fed with pure water, such that the minimum water usage is obtained through summing the flow rates through all the units. It is necessary to stipulate, for all units, the maximum inlet and outlet concentrations for the different contaminants of
A contaminant, in this context, is defined as any property of the water that prevents its direct reuse, and can thus include heat content (i.e. temperature) as well as the usual physicochemical attributes such as suspended solids, acidity and hardness. Increasing the allowable influent concentration results, in general, in an increased effluent concentration. In most cases basic rules can be applied to determine the maximum allowable effluent concentration and, as such, the maximum allowable influent concentrations, based around such fundamental properties as mass transfer, solubility of scalants, corrosivity and (organic) fouling. This relaxing of concentrations allows flow rates to be determined that are most appropriate for efficient water use.

*The limiting water flow rate concept (limiting water line)*

An important difference can be observed between two basic precepts concerning flow rate. The flow rate may either be fixed at some value or can be assumed to take on some limiting value whereby the maximum allowable inlet and outlet concentrations are obtained according assumptions or measurements made based on deterioration of water quality through a unit. In Fig. 4.9 these approaches are presented for a simple single-contaminant case. The physicochemical properties of the process and of the equipment allow a maximum inlet and effluent concentration of 50 ppm and 150 ppm respectively. However, when increasing the inlet concentration up to 50 ppm, and keeping to the existing flow rate of 2 t/h, the maximum effluent concentration is not reached for this fixed flow rate. Permitting a variable flow rate for the process considered allows the maximum effluent concentration to be reached. The water pinch methodology was initially presented as a problem without flow rate

![Figure 4.9](image-url)
constraints (Wang and Smith, 1994a) and later extended to fixed flow rate (Wang and Smith, 1995) since this situation is most common in process industries. However, the original methodology can be extended to processes with flow rate constraints.

The limiting composite line

Consider an example scheme with three processes (P1 to P3) and two contaminants (C1 and C2). It is assumed, for all processes, that the limiting flow rates and the initial flow rates are the same for the relaxed case, and that the flows have been optimised. Data for the example are provided in Table 4.4.

The pinch methodology begins with the construction of the limiting composite line. To this purpose separate limiting water profiles (solid, fine lines) are plotted from the data in Table 4.4 as a concentration versus mass load diagrams for both contaminants (Figs. 4.10 and 4.11). From Fig. 4.10 four concentration intervals can be distinguished (0–25; 25–50; 50–100; 100–150 mg/l) for C1, whereas only two intervals can be distinguished for C2 (Fig. 4.11) at 0–50 and 100–150 mg/l. In each concentration interval, a line is then constructed between the point at lowest mass load and concentration and the highest, such that limiting composite curves are obtained for both contaminants (solid, thick lines). For both contaminants the minimum flow rate through the overall scheme can now be determined by drawing a line that at no point bisects the limiting composite line. This line is called the water supply line, the inverse of the slope being the overall limiting flow rate. In this case, for both contaminants the water supply line only touches the composite curve at the end point. This point called the pinch point.

From Figs. 4.10 and 4.11 it is observed that two limiting flow rates are obtained: 2.67 t/h for C1 and 2.33 t/h for C2. In general, the theory of water pinch is presented in articles and textbooks as a single-contaminant case. Indeed, it is not possible to consider more than one contaminant since concentration shifting, as proposed by Wang and Smith (1994a) is required when targeting a multiple contaminant case. This entails a very lengthy and complex procedure. Although a two-contaminant case has been assumed to illustrate the method

### Table 4.4 Water data for example 1 after relaxation of the contaminants

<table>
<thead>
<tr>
<th>Process (P)</th>
<th>Mass load (g/h)</th>
<th>Concentration (ppm)</th>
<th>Limiting flow rate (t/h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Process 1 (P1)</td>
<td>100</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Process 2 (P2)</td>
<td>100</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>Process 3 (P3)</td>
<td>200</td>
<td>100</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Process (P)</th>
<th>Mass load (g/h)</th>
<th>Concentration (ppm)</th>
<th>Limiting flow rate (t/h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Process 1 (P1)</td>
<td>100</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Process 2 (P2)</td>
<td>50</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>Process 3 (P3)</td>
<td>200</td>
<td>100</td>
<td>4</td>
</tr>
</tbody>
</table>
and its application, nearly all practical cases concern multiple-contaminant matrices. It should be stressed that, although applicable for this case, it is not generally the case that the overall target flow rate, based on all contaminants in a multiple-contaminant system, corresponds to the highest value flow rate observed when constructing the water supply lines.
4.2.4 Computed solutions

For the simple scenario given in Table 4.4, implementation of both LM and UMIST software provides the same limiting flow rate, corresponding to the theoretical limit. However, the networks proposed by the two different software packages differ (Figs. 4.12 and 4.13). Indeed, by allowing a variable flow rate through the process, the UMIST software Water, which allows both approaches (i.e. fixed and non-fixed flow rates) projects flow rates lower than the limiting ones. Using the LM WaterTarget™ software, based on flow rates fixed at the limiting value, the individual flow rates in the processes remain the same. To achieve the same target flow rate, but constraining to a fixed process flow rate, WaterTarget™ proposes a network where part of the effluent of Process 2 and 3 is recycled and used as influent for those processes (Fig. 4.12). As such the overall flow rates through all the individual processes remains constant. The network proposed by Water looks the same as the WaterTarget™ one, although providing no recycling and thus allowing a reduced flow rate through the third process.

This example shows that whether or not the flow rate through the process is constrained at a fixed value, the target minimum flow rate remains the same and both software tools provide a possible water network achieving the target. Moreover, when the results of the optimisation indicate recycling of the outlet back to the inlet of the process, this suggests the possibility of lowering the flow rate through the process.

Since the two approaches result in a different network, the inlet and outlet concentrations of the individual processes will also differ (Table 4.5). By not constraining the individual process flow rates, lower inlet concentrations from the Water solution are obtained compared to those from the fixed flow rate approach of WaterTarget™. However, since the final effluent is produced by Process 3, the effluent concentration of this process is the same for both approaches.

![Figure 4.12 Network proposed by WaterTarget™ to achieve the target for example 1](image)
Further decreasing the minimum flow rate

As shown above, the graphical presentation of the problem provides a method of determining the minimum flow rate, taking into account the prevailing constraints. The task of any engineer dealing with water saving must now be to search for ways to further reduce the water consumption. The graphical presentation provides an aid in attaining this goal. (The graphical plot used in WaterTarget does not represent concentration versus mass load, but concentration versus flow rate. This methodology and the interpretation of the curves generated can be found in Buehner and Rossiter (1996).)

A decrease in the overall water usage implies an increase in the slope of the water supply line. To achieve this for a single-contaminant scenario it can be observed from Fig. 4.10 that the maximum effluent concentration of Process 3 should be allowed to increase since, provided the inlet concentration is unchanged, the slope of the limiting water line for this process will increase when not constraining the flow rate. As such, the slope of the dotted line, representing the water supply line, will increase resulting in a lower target. For the example presented, however, the gain is only minor. Indeed, one
immediately observes that the slope of the limiting composite curve can only increase to the point at which the line touches the composite curve at the point (125, 50). As such, the effluent concentration can only be allowed to increase up to 160 ppm, resulting in an overall target of 2.5 t/h. The limiting composite curve for the new problem is presented along with the water supply line and the pinch point in Fig. 4.14. It is observed that further increasing the effluent concentration of Process 3 provides no further advantage.

Whereas Water provides a graphical presentation of the composite curves, thus allowing the engineer to tackle the problem graphically, WaterTarget provides a visual representation. Performing a so-called inlet sensitivity analysis, WaterTarget pinpoints those processes and contaminants that may be adjusted so as to achieve an improved target. The result of this sensitivity analysis, which essentially corresponds to the graphical analysis described previously, is presented as a bar chart of cost saving in $/h per unit ppm concentration change for each contaminant (Fig. 4.15). For the example given in this section, the software advises the engineer that it is beneficial to increase the inlet concentration of Contaminant 1 in Process 3. For this simple case, the bar chart is perhaps not necessary. For a more complex case, however, it provides a more accessible illustration of the system data. Indeed, whilst the theory is only valid for a single-contaminant problem, the sensitivity analysis applies to multiple-contaminant cases.

As already observed, the different approaches of fixed and non-fixed flow rates provide different solutions. To permit an increased effluent concentration from Process 3 and maintain a constant flow rate through the process, the inlet concentration must be allowed to increase for the fixed flow rate approach. If

![Figure 4.14](image)

*Figure 4.14 Construction of water supply line for Contaminant 1 after relaxing the effluent concentration for Process 3*
increasing this influent concentration is not permitted, no further reduction of the flow rate is possible. However, as shown above, the difference between the results obtained using Water and WaterTarget™ concern only the stream recycle presented in the solution provided by WaterTarget™. Since this recycle is an indication to the engineer that lowering the overall flow rate through the process is beneficial and would lower the inlet concentration to the process, a higher influent concentration could be entered and the recycled stream dispensed with. This would lower the flow rate through one or more of the processes whilst still providing a solution that complies with the original constraints. The LM software does not provide a means of helping the engineer to tackle this problem, but experienced designers may choose to use this option.

From Fig. 4.14 it is evident that further decreasing the overall water usage is only possible by shifting the pinch point to higher concentrations. To achieve this, the only possible solution is to increase the inlet concentration of Process 2. Indeed, changing the parameters of Process 1 or 3 will not alter the position of the pinch point. In general processes above the pinch point should not be considered since they do not influence it at all; for processes below the pinch point the system should be analysed to identify which process or processes have a positive influence on the overall water demand.

To further develop the methodology for a multiple contaminant case, the solution provided by WaterTarget™ (where both inlet and effluent concentrations from Process 3 are adjusted to maintain the constant flow rate) is further developed. It is assumed that after careful examination the effluent concentration is allowed to increase up to 190 ppm, and thus the influent concentration forced to increase by the same amount to 140 ppm. The bar chart (Fig. 4.16) now indicates that the inlet concentration of C1 of Process 2 should be altered. (The representation may appear somewhat confusing if one is not well

**Figure 4.15** Sensitivity analysis graphically presented by WaterTarget™
acquainted with the software: it might be deduced that altering the inlet concentration of C2 to Process 1 is more beneficial, but the negative sensitivity for C1 in the same process provides an indication that this option is not beneficial. Indeed, changing C2 has no influence in this case.) The bar chart representation for the inlet sensitivity accounts for both the inlet and outlet concentrations. The negative concentration indicated in the figure implies that it is beneficial to lower the inlet concentration, which results from an analysis of the outlet concentration. Since the inlet and outlet are coupled the software advises a change in inlet concentration since this is more practical.

In allowing the inlet concentration of C1 of Process 2 to increase up to 37.5 ppm and the effluent concentration to 137.5 ppm, and concomitantly allowing C1 in Process 3 to increase up to 200 ppm, WaterTarget indicates a target of 2.33 t/h. This value can, in this case, easily be explained using the graphical presentation of the UMIST software. Indeed, according to Fig. 4.17 a target of 2.25 t/h can be achieved for C1. However, this is not the overall target as the target set for C2 indicates a flow rate of 2.33 t/h, slightly higher than the one for C1. Therefore the overall target is 2.33 t/h. The process of identifying the minimum flow rate proceeds through an investigation of C2 (Fig. 4.18), and continues until no further adjustments to inlet and outlet concentrations can usefully be made.

In Figs. 4.19 and 4.20 water networks are presented for the WaterTarget and Water models respectively. The fixed flow rate approach of WaterTarget again produces a network with internal recycles, whereas Water proposes a lower flow rate through the process. As stated above, the net result for the latter is a lower inlet concentration to the processes (Table 4.6) with, in the case of Process 3, a C1 value which is below the original value of 100 ppm compared with a value of 121 ppm obtained from the WaterTarget analysis. One is
Figure 4.17  Water supply line for C1 after relaxing the effluent concentration for P3 and the influent and effluent concentrations for P2

Figure 4.18  Bar chart after relaxation for C1 in Processes 2 and 3

therefore confronted with the options of a change in either inlet concentration or flow rate.

It is important to stress at this point that water pinch provides a systematic approach for modelling an existing water network, rather than a single solution to a problem. The available software provides a useful tool for the engineer in the decision-making process, but identifying the actual minimum flow rate is an
iterative process. In the following paragraph an example is presented where it is assumed these first steps to determine the maximum inlet and outlet concentration have been fully examined.

4.2.5 Software tools currently available

As is already evident, application of water pinch methodology is simple for a single-contaminant process scheme and/or only water usage is to be considered without taking into account possible purification processes, discharge limits, or costs linked to water management and purification. Pinch methodology was originally developed as a research tool to help develop different scenarios for minimising water consumption for industrial case studies. With the expansion of water pinch as a design aid, commercial software packages have been developed.

Currently, there are at least four commercial software packages available: WaterTarget™ commercialised by Linnhoff March a division of KBC Process
Table 4.6 Resulting inlet and outlet concentration for the processes after relaxing C1 for P2 and P3

<table>
<thead>
<tr>
<th>Contaminant 1</th>
<th>Contaminant 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C1_in (ppm)</td>
</tr>
<tr>
<td>WaterTarget(\text{\textsuperscript{\textregistered}})</td>
<td></td>
</tr>
<tr>
<td>Process 1 (P1)</td>
<td>0</td>
</tr>
<tr>
<td>Process 2 (P2)</td>
<td>34</td>
</tr>
<tr>
<td>Process 3 (P3)</td>
<td>121</td>
</tr>
<tr>
<td>Water(\text{\textsuperscript{\textregistered}})</td>
<td></td>
</tr>
<tr>
<td>Process 1 (P1)</td>
<td>0</td>
</tr>
<tr>
<td>Process 2 (P2)</td>
<td>33</td>
</tr>
<tr>
<td>Process 3 (P3)</td>
<td>86</td>
</tr>
</tbody>
</table>

Technology limited (further referred to as LM software), Aspen Water\(\text{\textsuperscript{\textregistered}}\) commercialised by Aspen Technology, Water provided by UMIST to the members of its Research Consortium and Water Close\(\text{\textsuperscript{\textregistered}}\) commercialised by American Process with an emphasis on mills (American Process Inc. 2002; Retsina and Rouzinou, 2002).

Aspen Water\(\text{\textsuperscript{\textregistered}}\) supports the work flow at each stage of the design, addressing the total problem including process, utility and treatment systems. The company is able to call upon its library of standard water-using unit operation including boilers, evaporators, cooling towers, vessels, reactors and heat exchangers, as well as purification technologies such as membrane processes. The flow-sheeting structure allows users to develop new unit operation models for their own specialised application. Moreover, using Aspen Water\(\text{\textsuperscript{\textregistered}}\) allows the water chemistry to be modelled using the Aspen Properties\(\text{\textsuperscript{\textregistered}}\) physical property system. Additionally, using the Aspen Equation Modeler\(\text{\textsuperscript{\textregistered}}\), dynamic simulation is possible. Lourens (2002) provides an example of an industrial study performed using Aspen Water\(\text{\textsuperscript{\textregistered}}\).

The capability of WaterTarget\(\text{\textsuperscript{\textregistered}}\) is comparable to that of Aspen Water\(\text{\textsuperscript{\textregistered}}\) but it does not have the functionalities of a library containing different unit operations and treatment techniques. Simplified linear equations are provided to calculate the effluent concentration in unit operations based on the influent concentration and simplified treatment units are provided to mimic the processes involved. It is assumed the effluent concentration is inferred from the influent concentration and a single removal ratio determined for a specific contaminant. Cost functions are linearly dependent on the flow rate.

The project builder of WaterTarget\(\text{\textsuperscript{\textregistered}}\) (Fig. 4.21) allows different contaminants to be selected, preferably limited to five in number, the processes to be defined, the sources (mains water, ground water, etc.) and sinks (sewer, surface water, etc.) to be identified and treatment techniques to be selected. Any constraints on the system design can be imposed via the Bounds editor (Fig. 4.22). Several types of bounds may be inserted, for example restriction on flow rates, costs of connection of one process to another, or any
other logistical limitation. The water network obtained with the software is presented by means of a table, indicating all selected connections between sinks and sources. A visual presentation of the water network is not provided in WaterTarget\textsuperscript{\textregistered}, unlike Aspen Water\textsuperscript{\textregistered} and Water. Water, although being a research tool, does provide a graphical presentation of the network, thus allowing a very quick and easy assessment of the effect of changing constraints of the design.

Although the methodology for energy pinch was first extended by the UMIST Process Integration group to water systems, the software developed by this group has not really developed far beyond the research tool stage, although the Water software is in a continual state of development with new features being added regularly. Against this, Aspen Technology and Linnhoff March both entered the market in 1999 with more user-friendly software than that of the UMIST group, both groups having benefited from membership of the UMIST research group.
consortium which provides the latest information on the latest developments regarding the methodology itself.

Apart from the commercial software tools, several researchers have developed their own software tools, mainly from the mid-1990s onwards. This has arisen as much out of financial prudence as much as anything else, since commercial packages are both expensive and limited in that they do not allow the algorithms to be adapted. The simulation tool Matlab has been used by the Pollution Research Group at the University of Natal to formulate the problem and to design the water network in a power plant (Brouckaert et al., 2002). This formulation has been adapted by Majozi (1999) for batch processes. The group is also using this approach to optimise reagent recovery and water use in industrial processes. The software tool developed by the group to this purpose has yet to be commercialised (Giannada et al., 2002).

At the Technical University Hamburg-Harburg a water pinch software package, Wado\textsuperscript{w}, was developed with emphasis on the treatment system including the selection of reasonable treatment operations for purification (or regeneration) purposes (Ullmer et al., 2002). A software package (WaterDesign) for calculating minimum water usage for a single contaminant problem was provided along with a textbook (Mann and Liu, 1999). A Portuguese research group has developed its own software in using Microsoft Visual Basic, implementing the methodology of Wang and Smith (1994) with improvements including multiple pinch analysis, regeneration reuse targeting, taking into account water purification technology (Castro et al., 1999). According to the authors' article, the software is continually under development. At the University of Oklahoma the pinch concept was extended to account for heat and mass exchanger networks (Bagajewicz, 2002). Many other research groups and independent research institutions, such as CEVI in Denmark (Anderson et al., 2002), have produced their own CAD packages, and it is most likely that at least some of these will be commercialised at some stage.
Programming

The solution of a water demand problem based on pinch methodology essentially has two components: a proper routine that formulates the problem and accounts for the superstructure and a robust optimisation package to solve the highly non-linear equations. Optimisation demands the solution of highly non-linear equations. Indeed, the pinch problem is fundamentally non-linear with discrete decisions (complexity, number of connections, etc.), generally demanding solution by MINLP algorithms (Mixed-Integer Non-Linear Programming). For WaterTarget it can be observed that in running the optimisation procedure, GAMS (General Algebraic Modeling System; GAMS, 2002) is called upon as a solver. The commercial solvers used are OSL (MILP) and MINOS (NLP), whereas the algorithm used to solve the MINLP is proprietary. As for Water, the problem is decomposed into a linear approximation (MILP) followed by a non-linear (NLP) solver. Gianadda et al. (2002) use the UMIST approach, breaking down the problem into a MILP followed by NLP using CONOPT as the solver to deal with the NLP problem. The optimisation problem has been solved by Ullmer et al. (2002) by using a MINLP algorithm developed by Lassahn and Gruhn (2002). XPRESS-MP solves the MILP problem; the remaining NLP problem is solved by LSGRG2. At the Rzeszów University of Technology (Poland) an adaptive random search (ARS) optimisation method was developed to design the optimal wastewater reuse network containing a small or medium number of water-using processes (Poplewski et al., 2002). To overcome the limited number of processes, further investigations are planned. Shafiei et al. (2002) propose an alternative approach by coupling genetic algorithms with mathematical programming and evaluating the solutions obtained by process simulation.

Solving the highly non-linear problem forces the user, and thus the software developer, to carefully examine the solution to ensure that the solution provided is not a local but a global minimum. However, due to the non-linearity there is no guarantee the solution is a global one. Using many different initial conditions appropriate to the system being examined and searching the optimum value is an indication of how well the global optimum has been reached. The process of thoroughly checking this often makes the calculation lengthy, especially when many contaminants, regeneration techniques and constraints are considered. The software should indicate whether a local, a presumed global or no optimum has been reached.

4.2.6 Case study: water pinch and implementation of regeneration techniques

As already stated, water pinch is a methodology to determine the minimum water usage as an overall target and, as such, does not take into account the minimum total cost and compliance with effluent discharge consents. A case study is presented where a water network is determined, achieving minimum water usage at minimum total cost. The example is based on the WaterTarget software package, this being a package that allows a total system design to be developed, including compliance with the final effluent discharge limits. The theory behind the methodology incorporating regeneration techniques and final
effluent treatment can be found in El-Halwagi (1992) and Wang et al. (1994b). The case study illustrates the importance of software tools, comprising the water pinch methodology, to provide ideas promoting an integrated approach. It will be shown that, using available data from different purification techniques, the software package reveals the most cost-effective selection of segregated purification techniques and end-of-pipe treatment both to comply with the stringent effluent standards and to minimise the overall water consumption. The program allows the insertion of as many purification techniques as is desired, since it is not known in advance which purification technologies or the combination thereof provide the greatest cost benefit.

The textile process
Textile processing has been discussed in Section 2.3. It consists of many different steps, nearly all of them generating wastewater. The amount and composition of these individual effluent streams depends on many different factors, including the processed fabric and the process used (Bisschops, 2002). Due to more stringent effluent constraints and the increasing cost of water usage, much effort has been devoted to minimising water usage and improving product recovery in the textile industry.

A general flowchart of a textile manufacturing process comprises (Bisschops, 2002):

- **Desizing** with high BOD, COD and SS effluent concentrations,
- **Scouring/washing** with high BOD, COD, SS effluent concentrations,
- **Bleaching** with low BOD, high COD and solids effluent concentrations,
- **Singeing** where only cooling water is needed,
- **Cotton mercerising** with low BOD and SS effluent concentrations,
- **Dyeing** with very high COD and high SS effluent concentrations, with BOD levels dependent upon dye biodegradability,
- **Printing** with very high COD and high SS effluent concentrations and BOD levels dependent upon dye biodegradability, and
- **Finishing** where waters vary widely according to the type of finishing process and information regarding effluent concentrations and flow rates are scarce.

Apart from the water usage in these different process steps, water is used in the process for washing after pretreatment steps, after dyeing and after some finishing steps. Water is used for utilities such as raising steam, general cooling and maintenance.

Process data
The pure water source available for the processes is groundwater having the following specification with respect to six key contaminants:

- 5 ppm SS,
- 10 Pt-Co colour.
- 5 ppm BOD₅,
- 10 ppm COD,
- 0 ppm salt, and
- temperature of 12°C.

The cost price of this water is set at €0.1 per m³.

Flow rates and effluent concentrations used are listed in Table 4.7. The process layout has been simplified and, although most textile manufacturing processes are operated batchwise, the process is presented as a continuous one. The implications that need to be taken into account to transpose the results to the true batchwise operation are elaborated later.

For most processes the influent water quality has to be very high (equivalent to groundwater quality) since substances present in the process water can have an adverse impact on final product quality. Unfortunately, the exact requirements for wastewater reuse are not very clear and very few publication are available (Bisschops, 2002). For the purpose of the case study it is assumed that prior evaluation and discussion with the plant engineers, have established the maximum inlet concentrations and that no further relaxation is possible. The values used for the purpose of this example are shown in Table 4.8.

Wastewater data

The maximum values for discharge to the sewer system are set at: 100 ppm SS, 200 ppm BOD, 800 ppm COD, 1500 Pt-Co for colour and 30°C, with no limits for salt. The limits for discharge on the surface water are set at: 60 ppm SS, 25 ppm BOD, 125 ppm COD, 100 Pt-Co for colour and 30°C, again with no limits set for salt.

Table 4.7  Flow rates and effluent concentration for the processes in textile manufacturing (Bisschops, 2002; and others)

<table>
<thead>
<tr>
<th>Process</th>
<th>Flow (t/d)</th>
<th>SS (ppm)</th>
<th>Colour (Pt-Co)</th>
<th>BOD (ppm)</th>
<th>COD (ppm)</th>
<th>Salt</th>
<th>Temp. °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steam production</td>
<td>100</td>
<td>20</td>
<td>50</td>
<td>5</td>
<td>30</td>
<td>5</td>
<td>30</td>
</tr>
<tr>
<td>Cooling</td>
<td>350</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>40</td>
</tr>
<tr>
<td>Desizing</td>
<td>100</td>
<td>100</td>
<td>200</td>
<td>2700</td>
<td>11000</td>
<td>2</td>
<td>35</td>
</tr>
<tr>
<td>Prewashing</td>
<td>150</td>
<td>100</td>
<td>100</td>
<td>50</td>
<td>200</td>
<td>1</td>
<td>35</td>
</tr>
<tr>
<td>Bleaching</td>
<td>50</td>
<td>7900</td>
<td>500</td>
<td>200</td>
<td>8500</td>
<td>3</td>
<td>35</td>
</tr>
<tr>
<td>Mercerising</td>
<td>50</td>
<td>1250</td>
<td>200</td>
<td>55</td>
<td>100</td>
<td>1</td>
<td>35</td>
</tr>
<tr>
<td>Dyeing</td>
<td>300</td>
<td>190</td>
<td>4750</td>
<td>1400</td>
<td>1600</td>
<td>4</td>
<td>35</td>
</tr>
<tr>
<td>Printing</td>
<td>60</td>
<td>9500</td>
<td>10000</td>
<td>5000</td>
<td>50000</td>
<td>4</td>
<td>20</td>
</tr>
<tr>
<td>Final hot washing</td>
<td>50</td>
<td>50</td>
<td>200</td>
<td>20</td>
<td>50</td>
<td>0</td>
<td>35</td>
</tr>
<tr>
<td>Final cold washing</td>
<td>200</td>
<td>50</td>
<td>200</td>
<td>20</td>
<td>50</td>
<td>0</td>
<td>15</td>
</tr>
<tr>
<td>Maintenance</td>
<td>150</td>
<td>100</td>
<td>150</td>
<td>20</td>
<td>100</td>
<td>1</td>
<td>35</td>
</tr>
<tr>
<td>Total</td>
<td>1560</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*a It is assumed that cooling of the process baths lowers effluent temperature to 45°C. Due to a lag time in the installation before reaching the central wastewater treatment plant, the effluent temperature is taken at 35°C.
Table 4.8 Maximum influent concentration for the processes in textile manufacturing

<table>
<thead>
<tr>
<th>Process</th>
<th>SS (ppm)</th>
<th>Colour (Pt-Co)</th>
<th>BOD (ppm)</th>
<th>COD (ppm)</th>
<th>Salt (ppm)</th>
<th>Temp. (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steam production</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>25</td>
</tr>
<tr>
<td>Cooling(^a)</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>12</td>
</tr>
<tr>
<td>Desizing</td>
<td>5</td>
<td>100</td>
<td>50</td>
<td>200</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>Prewashing</td>
<td>50</td>
<td>2000</td>
<td>100</td>
<td>400</td>
<td>2</td>
<td>40</td>
</tr>
<tr>
<td>Bleaching</td>
<td>5</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>Mercerising</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>Dyeing</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>Printing</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>Final hot washing</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>1</td>
<td>60</td>
</tr>
<tr>
<td>Final cold washing</td>
<td>5</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>1</td>
<td>25</td>
</tr>
<tr>
<td>Maintenance</td>
<td>100</td>
<td>2500</td>
<td>200</td>
<td>400</td>
<td>4</td>
<td>60</td>
</tr>
</tbody>
</table>

\(^a\) The influent concentrations for cooling are set this strict to allow for later reuse of the resulting hot water for all steps in the manufacturing process.

Water discharged to the sewer system is set at a cost of €1 per m\(^3\), and to surface water (on meeting discharge consents) cost of €0.1 per m\(^3\). In reality the levies for discharge both on the sewer system and the surface water depend on the pollutant load in the effluent. Such a complex relationship cannot, however, be used in the software and instead fixed values must be used. Recalculation after optimisation is necessary to properly account for this.

The concentrated stream from the central wastewater treatment plant and the concentrated polluted streams from individual purification techniques can be treated separately in dedicated treatment facilities outside the textile company at a cost price of €100 per m\(^3\).

Purification techniques

Amongst the purification techniques (Schönberger, 1998) two important groups can be distinguished:

- processes capable of recovering of one or more of the reagents for reuse in the manufacturing processes, many of them also producing a highly purified effluent water which can easily be reused, and
- processes capable of recovering water for reuse purposes.

Data are summarised in Tables 4.9 and 4.10.

Selected purification techniques and cost data for these comprise the following.

Ultrafiltration for recovery of sizing agent (European Commission, 2001). The process is assumed to recover both the sizing agent, which can be reused in the process, and good-quality effluent water for reuse. Capital cost for a 100 m\(^3\) day\(^{-1}\) plant is around €1.2 million, including the UF plant, an equalisation tank, start-up and miscellaneous. Eventually a prefiltration step has to be accounted for. Savings up to €0.7 million per year due to product recovery.
Table 4.9 Removal efficiency (%) for the proposed purification techniques

<table>
<thead>
<tr>
<th>Method</th>
<th>SS (%)</th>
<th>Colour (%)</th>
<th>BOD (%)</th>
<th>COD (%)</th>
<th>Salt (%)</th>
<th>T (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>UF for recovery of sizing agent</td>
<td>100</td>
<td>75</td>
<td>95</td>
<td>90</td>
<td>95</td>
<td>Cst</td>
</tr>
<tr>
<td>Filtration after prewashing</td>
<td>65</td>
<td>0</td>
<td>40</td>
<td>50</td>
<td>50</td>
<td>Cst</td>
</tr>
<tr>
<td>Three-stage evaporation for recovery of alkali from mercerising</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>50</td>
</tr>
<tr>
<td>Membrane treatment for recovery of wastewater from printing paste</td>
<td>100</td>
<td>95</td>
<td>95</td>
<td>90</td>
<td>85</td>
<td>Cst</td>
</tr>
<tr>
<td>Membrane treatment for recovery of wastewater from final washing</td>
<td>95</td>
<td>60</td>
<td>85</td>
<td>90</td>
<td>85</td>
<td>Cst</td>
</tr>
<tr>
<td>Centralised biological WWTP</td>
<td>99</td>
<td>90</td>
<td>100</td>
<td>95</td>
<td>10</td>
<td>15</td>
</tr>
<tr>
<td>RO for the treated wastewater</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>Cst</td>
</tr>
</tbody>
</table>

*Removal efficiency by the centralised WWTP will largely depend on the final effluent quality for treatment. If, for example, the printing paste is to be treated in the WWTP as well, COD removal will be much lower. These water-insoluble compounds are substantially undegraded biologically, and are removed only by adsorption onto the sludge. Such removal and other special cases are not accounted for in this study.

Table 4.10 Water recovery ratio for the proposed purification techniques

<table>
<thead>
<tr>
<th>Method</th>
<th>Water recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>UF for recovery of sizing agent</td>
<td>87</td>
</tr>
<tr>
<td>Filtration after prewashing</td>
<td>97</td>
</tr>
<tr>
<td>Evaporation for recovery of alkali from mercerising</td>
<td>97</td>
</tr>
<tr>
<td>Membrane for recovery of wastewater from printing paste</td>
<td>90</td>
</tr>
<tr>
<td>Membrane for recovery of wastewater from final washing</td>
<td>95</td>
</tr>
<tr>
<td>Centralised biological WWTP</td>
<td>90</td>
</tr>
<tr>
<td>RO for the treated wastewater</td>
<td>85</td>
</tr>
</tbody>
</table>

alone can be achieved. Therefore the capital cost is assumed to be entirely recovered by the sizing agent recovery. Operating cost is assumed to be €0.1 per m$^3$, the figure being substantially reduced by the economic savings from the product recovery. It is assumed that a reusable effluent is provided at the same price as that of groundwater.

Filtration after prewashing. Capital cost is assumed to be €0.1 per m$^3$ (10-year depreciation period at an interest rate of 0%) for a self-cleaning rotary filter or a pressurised microfiltration membrane with a capacity of 150 m$^3$ day$^{-1}$. Operating cost is assumed to be €0.1 per m$^3$.

Three-stage evaporation for recovery of alkali from mercerising (European Commission, 2001). The process produces a concentrated lye (alkali) which can be reused in the process, and also provides a good quality effluent water for reuse. Capital cost depends on the plant size and the purification technique applied after evaporation to obtain a concentrated lye, which can be reused in the process, and typically varies between €0.2 million and €0.8 million. Due to product lye recovery the payback time can be less than one year, and the capital cost is therefore ignored in this study as it was for the ultrafiltration unit for recovery of
sizing agent. Operating cost is assumed to be €0.1 per m³, based on the same assumption of a cost recovery as the IJF plant.

Membrane treatment for recovery of wastewater from printing paste (European Commission, 2001). Capital cost for a 60 m³ day⁻¹ plant is €0.82 per m³ based on a 10-year depreciation period at an interest rate of 0%. The installation comprises coagulation/precipitation pretreatment followed by a microfiltration unit using a polypropylene membrane and a tube settler to remove the suspended solids. Operating cost, including external disposal of the concentrate, the principal contribution to opex, is about €4 per m³.

Membrane treatment for recovery of wastewater from final washing. Capital cost is assumed to be €0.33 per m³ for an ultrafiltration installation of 250 m³ day⁻¹. Operating cost is assumed to be €0.25 per m³.

Centralised biological wastewater treatment plant (European Commission, 2001). Capital cost is taken at €1.25 per m³ for a 2500 m³ day⁻¹ plant (10-year depreciation at an interest rate of 6.88%). The investment cost given in the paper (European Commission, 2001) was €1.50 per m³, although this included further purification (by reverse osmosis) of the effluent for reuse. The plant comprises equalisation, neutralisation, and activated sludge treatment in a special system of loop reactors and clarifiers, adsorption to remove dyestuffs, flocculation/precipitation and removal of sludge, and filtration in a fixed-bed gravel filter to remove suspended solids. Also included is the treatment of the sludge in a thickener and decanter and thermal regeneration in a rotary kiln. Operating cost is taken at €0.6 per m³, based on the above, including ash disposal and ignoring heat recovery.

Reverse osmosis for polishing the treated wastewater (European Commission, 2001). Capital cost is taken at €0.25 per m³ for a 1500 m³ day⁻¹ plant. Investment comprises the reverse osmosis plant consisting of 10 modules containing 4 spiral membrane elements each. Included is a storage tank conditioned with ozone to control biological growth. Operating cost is taken at €0.1 per m³.

Results

Base case: targeted minimum flow rate, discharge to sewer. For the base case it is assumed the company is required to meet stricter consents for discharge to sewer. In the first instance a large central wastewater treatment plant can be envisaged for the treatment of the overall wastewater.

The result of the base case reveals that 1210 m³ day⁻¹ of fresh water is needed with all effluent from the cooling process being recycled (350 m³ day⁻¹ waste from a 1560 m³ day⁻¹ feed). No other effluents are identified for reuse taking into account minimum operating cost. The effluent of the central wastewater treatment plant meets the strict discharge consents for both SS and BOD. Only 378 m³ day⁻¹ needs to be treated in the central plant, the remaining effluents being discharged directly to sewer. The daily cost of this base case is €6,387, 60% of this being due to the external treatment of the concentrated waste.

Scenario 1: targeted minimum flow rate, all purification techniques permitted. For this case it is presumed the company was seeking alternative ways of minimising the
overall cost. All purification techniques previously outlined are permitted and the pinch methodology is used to target the optimal solution.

As is already obvious from Section 4.2.4, recovery of the sizing agent and recovery of alkali with simultaneous recovery of a purified water is economically very beneficial, with water available at good quality at the cost price of groundwater. Looking at the proposed target (1016 m$^3$ day$^{-1}$ of freshwater), the water network suggested is operated at a cost price of €4819, lower than the base case. The target is achieved by implementing the membrane technique for treatment of the printing paste. The final effluent is discharged to the sewer system without implementing other purification processes to achieve a wastewater that can be discharged at a lower cost to the surface water. One of the flow rates suggested is very small and transpires to be an internal recycle for one process. The question of whether or not the overall water flow through the process should be lowered can then be addressed.

As can be seen in Table 4.11 for several processes multiple inlet streams should be connected. This often implies additional cost, not yet accounted for by the analysis, for pipework, storage tanks and control. It is advisable to ignore these costs when embarking on the analysis, since they make the problem too complex. When the optimum network is selected these additional costs can be added manually or can be taken into account in the program.

Scenario 3: targeted minimum flow rate when all treatments are allowed but discharge on surface water is imposed. The limits set for the discharge to surface waters are much stricter than those for discharge to sewer. When the company is faced with these stricter values, proper solutions have to be accounted for. Using the software for the case study and restricting discharges to surface water yields a daily operational cost of €6551. The solution indicates that the constraints can only be met when allowing good quality water, i.e. the outlet of the RO installation, to be discharged to the surface water. Indeed, the RO is essential for the strict regulations set to be met. It is observed that this good quality water is not being reused in the process. Careful examination of the problem reveals this is due to the low cost of the fresh water! Indeed, should the company be forced to cut down on the groundwater use, tap water with additional treatment to reduce the hardness, has to be used. A higher price of this water source would make the RO installation competitive.

Remarks

When critically evaluating the case study it is apparent that not all of the contaminants behave in a way that is consistent with necessary assumptions made in the pinch analyses. Indeed, it is assumed by the program that, for each contaminant, there is a linear relationship between the measurable concentration and the actual mass per unit volume when different streams are combined. For colour, this is not always valid. When running the program it is observed that, depending on the initial conditions, the software sometimes provides a solution that is indicated not to be the optimal one. Running the software again with different initial values can result in the optimal solution.
Table 4.11 Proposed water network to achieve the target set for Scenario 1

<table>
<thead>
<tr>
<th>From process...</th>
<th>To process</th>
<th>Flow (t/d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cooling out</td>
<td>Dyeing in</td>
<td>248.50</td>
</tr>
<tr>
<td>Cooling out</td>
<td>Final hot washing in</td>
<td>1.50</td>
</tr>
<tr>
<td>Cooling out</td>
<td>Desizing in</td>
<td>100.00</td>
</tr>
<tr>
<td>Prewashing out</td>
<td>Maintenance in</td>
<td>44.87</td>
</tr>
<tr>
<td>Dyeing out</td>
<td>Maintenance in</td>
<td>13.26</td>
</tr>
<tr>
<td>Final hot washing out</td>
<td>Prewashing in</td>
<td>50.00</td>
</tr>
<tr>
<td>Final cold washing out</td>
<td>Prewashing in</td>
<td>11.60</td>
</tr>
<tr>
<td>Maintenance out</td>
<td>Maintenance in</td>
<td></td>
</tr>
<tr>
<td>From utility...</td>
<td>...to process</td>
<td></td>
</tr>
<tr>
<td>Fresh water</td>
<td>Steam production in</td>
<td>100.00</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Cooling in</td>
<td>350.00</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Prewashing in</td>
<td>88.40</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Bleaching in</td>
<td>50.00</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Dyeing in</td>
<td>51.50</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Final cold cleaning in</td>
<td>200.00</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Maintenance cleaning in</td>
<td>65.76</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Mercerising in</td>
<td>50.00</td>
</tr>
<tr>
<td>Fresh water</td>
<td>Printing in</td>
<td>60.00</td>
</tr>
<tr>
<td>UF for size recover clean</td>
<td>Maintenance in</td>
<td>25.84</td>
</tr>
<tr>
<td>Evaporation for alkali recovery clean</td>
<td>Final hot washing in</td>
<td>48.50</td>
</tr>
<tr>
<td>From process...</td>
<td>...to utility</td>
<td></td>
</tr>
<tr>
<td>Steam production out</td>
<td>Discharge to sewer</td>
<td>100.00</td>
</tr>
<tr>
<td>Prewashing out</td>
<td>Discharge to sewer</td>
<td>105.13</td>
</tr>
<tr>
<td>Bleaching out</td>
<td>Discharge to sewer</td>
<td>3.09</td>
</tr>
<tr>
<td>Bleaching out</td>
<td>Centralised WWTP</td>
<td>46.91</td>
</tr>
<tr>
<td>Dyeing out</td>
<td>Discharge to sewer</td>
<td>97.09</td>
</tr>
<tr>
<td>Dyeing out</td>
<td>Centralised WWTP</td>
<td>189.65</td>
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<tr>
<td>Final cold cleaning out</td>
<td>Discharge to sewer</td>
<td>188.40</td>
</tr>
<tr>
<td>Maintenance out</td>
<td>Discharge to sewer</td>
<td>149.73</td>
</tr>
<tr>
<td>Desizing out</td>
<td>UF for size recovery inlet</td>
<td>100.00</td>
</tr>
<tr>
<td>Mercerising out</td>
<td>Evaporation for alkali recovery in</td>
<td>50.00</td>
</tr>
<tr>
<td>Printing out</td>
<td>Membrane for printing paste in</td>
<td>60.00</td>
</tr>
<tr>
<td>From utility...</td>
<td>...to utility</td>
<td></td>
</tr>
<tr>
<td>UF for size recover clean</td>
<td>Discharge to sewer</td>
<td>61.16</td>
</tr>
<tr>
<td>UF for size recover dirty</td>
<td>Recovered product</td>
<td>13.00</td>
</tr>
<tr>
<td>Evaporation for alkali recovery dirty</td>
<td>Recovered product</td>
<td>1.50</td>
</tr>
<tr>
<td>Membrane for printing paste clean</td>
<td>Discharge to sewer</td>
<td>54.00</td>
</tr>
<tr>
<td>Membrane for printing paste dirty</td>
<td>Centralised WWTP inlet</td>
<td>6.00</td>
</tr>
<tr>
<td>Centralised WWTP clean</td>
<td>Discharge on sewer</td>
<td>218.30</td>
</tr>
<tr>
<td>Centralised WWTP dirty</td>
<td>Dirty discharge</td>
<td>24.26</td>
</tr>
</tbody>
</table>

From the author's own observation, it appears to be more productive to start with initial conditions which are randomly selected, thus not necessarily close to the optimum.

The water pinch methodology described in Sections 4.2.1–4.2.4 was restricted to targeting the minimum flow rate reusing effluent streams without prior treatment. The methodology, however, also allows the identification of those processes for which treatment of the effluent prior to reuse should be
considered. For the case study, as presented above, it is assumed that the methodology is applicable. However, it is often beneficial, in practical cases, to insert additional purification methods in the software. The software then provides an accurate solution, and a decision must then be made as to whether it should be implemented.

Although the case study is limited to the textile process, it should be stressed that the pinch methodology applies across all industrial sectors and all plant sizes, from large power generation (Selby and Tvedt, 1998) and paper fabrication (Shafiei et al., 2002) plants to relatively small-scale operations within the pharmaceutical and food industries (Thevendiraraj et al., 2001). However, it is apparent both from examination of the literature and from personal contact with engineering firms and research institutes, that real values indicating targeted water usage and/or achieved water usage are very difficult to find, largely due to their commercial sensitivity. Moreover, it is very often difficult to be sure that the indicated result has actually been implemented. However, it can be stated that water savings of at least 20–60% are achievable through applying water pinch, as revealed from some of the references cited below.

4.2.7 Conclusion

Software is currently available, based on the water pinch methodology, that allows one to target the minimal water usage at minimal cost, taking into account different constraints. Although the methodology is simple when considering only one contaminant, no purification techniques and no costs, the methodology requires a skilled engineer or researcher to unravel the whole concept when taking into account all these elements. Fortunately, the software currently available provides a means for process engineers and researchers to tackle these problems without necessitating a fundamental understanding of the underlying concepts. The software provides a guide that helps the user through the different steps. Moreover, it is obvious that the software can easily be used to evaluate many different scenarios and investigate the influence of many parameters. However, the software tool should not be considered as a plug-and-play direct answer to the problem. Indeed, without proper insight into the methodology, processes might be overlooked that play an important role in reducing the overall water consumption. Identifying solutions that reduce water consumption at the lowest total cost demands the combined skills of both the process engineers of the problem holder and experts in water pinch methodology.

References


Majozi, T. (1999). The application of pinch technology as a strategic tool for rational management of water and effluent in an agrochemical industry. MSc ling dissertation, Pollution Research Group, School of Chemical Engineering, University of Natal, Durban, South Africa.


4.3 Design examples

4.3.1 Problem in reverse osmosis: film theory and energy demand

A maximum concentration polarisation parameter value of 1.14 is recommended for operation of a membrane element of the following specifications:

- Length \((L)\): 1 m
- Channel thickness \((h)\): 0.8 mm
- Membrane area \((A)\): 36 m²
- Spacer mesh width \((m)\): 0.6 mm

It can additionally be assumed that the ion diffusion coefficient is \(8 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}\), and the fluid viscosity and density values are \(1.15 \times 10^{-3} \text{ kg m}^{-1} \text{s}^{-1}\) and \(1000 \text{ kg m}^{-3}\) respectively.

If the element operates at a mean flux of 21 LMH, what is the minimum feed flow rate and what conversion does this yield?

If:

(a) the hydraulic losses amount to 1.15 bar per m/s cross-flow velocity per m path length.
(b) the membrane resistance is \(8.5 \times 10^{13}\) per m.
(c) the feedwater contains 850 mg l⁻¹ sodium chloride, and
(d) the membrane has a rejection of 98%, the water temperature is 15°C and \(y = 0.9\)

what feed pressure is required, and what energy demand does this equate to for a pumping efficiency of 40%?

Solution

Film theory states that the flux \(J\) and concentration polarisation parameter \(\varphi\) (i.e. \(c^*/c\)) are related by (Equation (2.14)):

\[ J = k \ln \varphi \]

The mass transfer coefficient is thus:

\[ k = \frac{J}{\ln \varphi} = \frac{21/(1000 \times 60 \times 60)}{\ln 1.14} = \frac{5.83 \times 10^{-6}}{0.131} = 4.45 \times 10^{-5} \text{ m/s} \]

The Sherwood number, according to Equation (2.15), is then:

\[ Sh = \frac{kd}{D} = \frac{4.45 \times 10^{-5} \times 1.6 \times 10^{-4}}{8 \times 10^{-10}} = 89.0 \]

since, for a parallel flow channel, \(d\) is equal to twice the channel thickness \(h\).
According to the expression derived for filled channels by Chiolle et al. (1978), Table 2.11 and Equation (2.15), the Sherwood number correlates to the Reynolds and Schmidt numbers $Re$ and $Sc$ respectively according to:

$$Sh = 1.065Re^{0.5}Sc^{0.33}[d/(6Lm)]^{0.5}$$

and from Equations (2.17) and (2.18):

$$Re = \frac{\rho Ud}{\mu} = 10^3 \times U \times 1.6 \times 10^{-3}/(1.15 \times 10^{-3}) = 1391U$$

$$Sc = \frac{\mu}{\rho D} = 1.15 \times 10^{-1}/(10^3 \times 8 \times 10^{-10}) = 1438$$

The Reynolds number can then be calculated as:

$$Re = \left( \frac{89}{1.065 \times 1438^{0.33}\left(\frac{1.6}{6 \times 1 \times 0.6}\right)^{0.5}} \right)^2 = 129$$

It follows that the (retentate) cross-flow approach velocity is:

$$U = \frac{129}{1391} = 0.0927 \text{ m/s}$$

The volumetric flow $Q$ at this flow velocity relates to the channel cross-sectional area, given by $d \times A/(2L)$:

$$Q = U \times d \times A/(2L) = 0.0927 \times 0.8 \times 10^{-3} \times (36/2 \times 1) \times 60 \times 60$$

$$= 4.81 \text{ m}^3/\text{h}$$

If this is assumed to relate to the flow at the retentate outlet, where concentration polarisation is greatest, then the feed flow will higher than this due to conversion. The total volumetric flow of permeate is given by:

$$Q_{\text{permeate}} = J \times \text{area} = (21/1000) \times 36 = 0.76 \text{ m}^3/\text{h}$$

The feed flow is then given by:

$$Q_{\text{feed}} = 4.81 + 0.76 - 5.57 \text{ m}^3/\text{h}$$

and the conversion is thus:

$$\Theta = 100 \times 0.76/5.57 = 13.6\%$$

This is within the normal range of 7–17% quoted for a typical brackish water 40-inch membrane element. However, the production rate is considerably lower.
than the flows of up to 1.6 m³/h attainable in practice for elements of this size, suggesting that mass transfer promotion is greater in real systems. This is partly accounted for by the higher cross-flows at the element inlet, but is most likely to be due to assumptions made about the respective values of the spacer mesh width and the diffusion coefficient. The calculation is extremely sensitive to both these values.

The feed pressure can be estimated from resistance theory, whereby:

- the hydraulic losses across the retentate side, and
- the transmembrane pressure

are calculated from the retentate channel hydraulic resistance and the membrane resistance \( R_m \) (Equation (2.5)). The feed pressure is further increased by the effect of concentration polarisation, since:

\[
\text{TMP} = J \times R_m \times \mu + \Delta \Pi^*.
\]

where

\[
JR_m\mu = 5.83 \times 10^{-6} \times 8.5 \times 10^{13} \times 1.15 \times 10^{-1} = 5.70 \times 10^5 \text{ Pa}
\]


dis - 5.7 bar

and

\[
\Delta \Pi^* = \varphi \Delta \Pi
\]

\[
\Delta \Pi = \gamma RT \Sigma \Delta C_i
\]

where \( \Delta \Pi \) is the maximum osmotic pressure difference across the membrane (from Equation (2.9)) modified for concentration polarisation and conversion. So:

\[
\Delta \Pi = 1.14 \times 0.9 \times 8.314 \times (273 - 15) \times 2 \times 0.98 \times (850/58.5)
\]

\[
= 69,960 \text{ Pa}
\]

Thus

\[
\Delta \Pi^* = 1.14 \times 69,960 = 7.98 \times 10^4 \text{ Pa}
\]

And so

\[
\frac{\Delta \Pi^*}{(1 - \theta)} = \frac{7.98 \times 10^4}{(1 - 0.136)} = 9.24 \times 10^4 \text{ Pa} = 0.924 \text{ bar}
\]

Therefore

\[
\text{TMP} = (5.70 + 0.92) = 6.62 \text{ bar}
\]
Now hydraulic losses across the retentate are given by
\[ \Delta p_{\text{losses}} = 1.15 \text{ UI bar for } U \text{ m/s and } L \text{ m} \]

where \( U \) is the mean retentate velocity
\[ U = \frac{0.0927}{2} \left( \frac{1}{1 - 0.136} + 1 \right) = 0.100 \text{ m/s} \]

So
\[ \Delta p_{\text{losses}} = 1.15 \times 0.100 \times 1 = 0.115 \text{ bar} \]

Thus, total pressure is given by \( \text{TMP} + \Delta p_{\text{loss}} = 6.74 \text{ bar} \).

According to Equation (2.23), the hydraulic energy demand per kg product is given by:
\[ E = \frac{(1/\rho \Theta) \Delta P}{60 \times 60} \]

Converting to kWh per m\(^3\) product:
\[ E = \frac{1000}{60 \times 60 \times 1000 \times \rho \Theta} \Delta P = \frac{2.78 \times 10^{-7}}{0.136} \Delta P \]
\[ = \frac{2.78 \times 10^{-7}}{0.136} \times 6.74 \times 10^5 = 1.38 \]

For a pumping efficiency of 40\%, this figure becomes 3.44 kWh m\(^{-3}\).

### 4.3.2 Problem in reverse osmosis: array design

If elements of the specification given in the previous problem are to be used to achieve an overall conversion of at least 75\% within an array, what array design can be used to deliver 35 l/s permeate product and what would be the specific energy demand?

**Solution**

For a conversion of 75\%, the feed and permeate flows are related by:
\[ Q_R = Q_F / 0.75 = 35 / 0.75 \text{ l/s} = 46.7 \text{ l/s} = 168 \text{ m}^3 / \text{h} \]

The overall conversion of 75\% suggests a 2:1 array, since conversion of 50\% at each stage produces 25\% conversion overall. The retentate flow from the module \( Q_R \) is then half of the feed flow \( Q_F \).

If the relationship given in Equation (2.3) is extended to a number of elements in series, then the retentate flow \( Q_R \) is related to the feed flow \( Q_F \) and the number of elements per module \( n \) by:
\[ Q_R = Q_F (1 - \Theta)^n \]

Hence:

\[ n = \frac{\log(Q_R/Q_F)}{\log(1 - \Theta)} \]

If a feed flow of 5.6 m³/h and a mean conversion of 13.6% per element is assumed then:

- Total number of rows = 168/5.6 = 30, and
- Elements per module = \log 0.5/\log 0.864 = 4.74 \sim 5.

Hence, an array comprising 30 rows of modules in the first stage and 15 in the second stage is needed, each module containing 5 elements. This would then give an overall recovery of:

\[ \Theta = 1 - (\Theta_{\text{overall}}^{n}) = 1 - 0.864^{10} = 76.8\% \]

Although 2:1 arrays are most common, it is also possible to employ a 3:2 array, which in this case would be 30:20:10. A third stage would be required to obtain the overall conversion of 75%, but the elements per module demanded to achieve 33% conversion for the first two stages would be less:

Elements per module, stages 1 and 2 = \log 0.667/\log 0.864 = 2.77 \sim 3

Stage 3 would then demand 4 elements per module to achieve the target overall recovery, since \( n_{\text{overall}} \) should still be 10 across the whole array.

A comparison of the two schemes (see below) reveals that the 2:1 array employs 15 fewer modules, but that the 3:2 array uses 35 fewer elements. Since the pressure vessels are expensive and extra pipework is demanded by having a third stage, the 2:1 array would usually be preferred on the basis of capital costs. On the other hand, there is a greater decrease in flow velocity across the module for the 2:1 array which inevitably leads to greater hydraulic loading at the front of the module and/or greater concentration polarisation at the back.

<table>
<thead>
<tr>
<th>Scheme</th>
<th>No. of modules</th>
<th>No. of elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 stages (2:1 array)</td>
<td>45</td>
<td>225</td>
</tr>
<tr>
<td>2 stages (3:2 array)</td>
<td>60</td>
<td>190</td>
</tr>
</tbody>
</table>

The specific energy demand relates to the total pressure and conversion, as before. The total pressure drop across the retentate is given by:

\[ \Delta P_{\text{retent}} = 1.15 \times U_{\text{mean}} \times (L \times n) \]
where \( n \) is the number of elements across the module. Estimation of the mean retentate flow velocity presents a problem. If the velocity at the outlet must be kept above \( 0.927 \, \text{m/s} \) to suppress CP then this implies that the module inlet velocity must be double this value, assuming 50% conversion, giving a mean velocity of \((3/2) \times 0.0927 = 0.139 \, \text{m/s}\). However, maintaining this velocity would necessitate an increase in the retentate flow through the module, which would then reduce the conversion unless this flow increase were to be produced by retentate recirculation. For the purposes of this calculation a mean retentate velocity of \( 0.139 \, \text{m/s} \) is assumed, notwithstanding the above implications.

\[
\Delta P_{\text{retent}} = 1.15 \times 0.139 \times (10 \times 1) = 1.60 \, \text{bar}
\]

The maximum osmotic pressure difference is given as before by:

\[
\frac{\Delta \Pi^*}{1 - \Theta} = \frac{1.14 \times 70000}{(1 - 0.768)} = 344 \, 000 \, \text{Pa} = 3.44 \, \text{bar}
\]

Thus, the total pressure, including the pressure drop due to the membrane itself, is:

\[
P = 5.70 + 1.60 + 3.44 = 10.7 \, \text{bar}
\]

and the specific energy consumption in kWh m \(^{-3}\) product is

\[
E = \frac{2.78 \times 10^{-7}}{\Theta} \frac{\Delta P}{0.768} = \frac{2.78 \times 10^{-7} \times 10.7 \times 10^{5}}{0.768} = 0.391
\]

For a pumping efficiency of 40%, this figure becomes 0.978 kWh m\(^{-3}\), considerably lower than the single element case because of the increased conversion and minimal extra energy expenditure.

### 4.3.3 Problem in reverse osmosis: CAD array design

Produce a complete array design based on the previous design specification for a feedwater of the following composition: 200 mg l\(^{-1}\) hardness "as CaCO\(_3\)", 150 mg l\(^{-1}\) alkalinity "as CaCO\(_3\)", 75 mg l\(^{-1}\) sulphate, \(\sim\)1200 mg l\(^{-1}\) total dissolved solids, pH 7.5, temperature 15°C, with an overall rejection of at least 98% and a product water pH of \(\sim\)7.0.

**Solution**

As already stated (Section 4.1), all of the leading RO manufacturers produce CAD packages which allow the designer to produce a complete array design for any specified combination of feed and product water quality. The key design parameters for any RO CAD package are as follows:
• feed or product water quality and quantity.
• membrane type, which then determines the flux and pressure, and
• conversion.

All of these parameters must be entered into the design before the array can be optimised, usually through iteration.

The comprehensiveness, level of sophistication, and user-friendliness of the design package varies from one manufacturer to the next. However, all these packages have been developed to help the designer produce an array design as easily as possible, using the manufacturer’s own RO membrane products. Hence the information pertaining to mass transfer of dissolved solids and water across the membrane is specific to the membrane products listed within the package. Apart from the way in which the design information is set out, this is the main difference between the available packages.

The ROPRO package (Version 7) is the proprietary CAD program produced by the suppliers of the Fluid Systems RO membranes. ROPRO offers a number of overall design configurations and options, such as bypass, pre- and post-acid/base dosing and degassing, and has a comprehensive “help” file. It also has an ancillary costing facility. Most of the ROPRO features are self-evident from the main flowsheet schematic, which has active regions for the pre-treatment, array and post-treatment processes, as well as for the stream compositions. Any of these can be checked at any point.

The quickest route to designing the plant is as follows:

1. Use the Input Wizard (on the Input menu) to enter all the data (ignoring pre-dosing for now)

2. Once the array design screen is reached, the Array Wizard should be used and an estimate of the flux made from the information given for the membrane element selected.

3. When all the data has been entered, the large GO button on the menu bar can be clicked to reveal the results.

The next screen informs the user where the design fails, either through figures shown in red for the scaling indices or through a Warnings prompt in the light blue box. A further screen appears when the Warnings button is pressed. Having read the warnings, the array design can be adjusted accordingly.

A few common remedial measures are as follows:

• RO membranes tend to scale-up on the concentrate side of the membrane. The only option offered by this package for ameliorating this problem is acid dosing.
• The flux chosen is critical. Too high a value causes an overlarge operating pressure and scaling problems. Too low a flux means that the system is hydraulically unbalanced and not optimised.
• The number of elements in each module can be adjusted, but the default value is normally quite reasonable.
Membranes for Industrial Wastewater Recovery and Re-use

The full details of the design may be viewed at any time from the File menu (Preview All) or by the short-cut key command Ctrl V.

For the problem specified it must be assumed that the hardness is all as calcium and the alkalinity as bicarbonate, so that these can be directly entered into the Feed Analysis screen. Composition values must be entered in the appropriate units (mg/l or mg/l as CaCO₃ in this case). An estimate of the sodium or chloride concentration is needed to complete the analysis and achieve the target TDS of 1200 mg/l. The package automatically adjusts the specification with the counter-ion (i.e. chloride or sodium ions) to obtain electroneutrality.

On the following pH Adjustment screen the required pH to which the water feeding into the array must be entered. It is simpler to do this once the scaling propensity (via the Langelier Scaling Index, LSI) has been calculated. On the following Flow Rates and Recovery menu the product flow rate (3024 m³/day and the recovery (75%) must be entered.

Next screen is the Membrane Array Configuration, which includes an Array Wizard that allows an estimate of the flux to be entered. RO membranes will normally operate at a flux of between 20 and 50 LMH, depending on the degree of fouling. Membrane elements can be selected from the left-hand drop-down menu and modules from the right hand one. For this large flow rate, a large membrane module is appropriate (prefix "8" for 8-inch diameter modules and MAG for the largest membrane element area). Having entered a flux value, the preliminary design is complete and the GO button can be pressed.

If a fairly conservative flux of 25 LMH is chosen for the array design, the only error arising will be from the scaling propensity of the water. This can be corrected by acid addition to being the LSI below zero. Again, trial and error is required for this, and through iteration it can be established that adjusting the pH to around 6.1–6.3 (depending on the membrane selected) by pre-dosing with either sulphuric or hydrochloric acid is sufficient.

The product water quality must then be checked to establish if 98% rejection has been achieved (i.e. the permeate product TDS is no more than 24 mg/l, 2% of the feed TDS of 1200 mg/l) and that the pH is around 7. The permeate product pH is always acidic due to the unrestricted permeation of carbon dioxide, which can be removed by stripping and/or neutralised by lime or caustic dosing. In reality, the controlling the product pH to 7 following CO₂ stripping would be extremely difficult for such a low TDS water. The permeate TDS can also, of course, be altered by membrane selection. Selection, once again, proceeds by trial and error – and more than one option is possible. A possible design is detailed below.

Pretreatment: Dosing to pH 6.2 with 93.2% sulphuric acid at 376.3 kg/day
Membrane: TFC 8832HR-575MAG
Array: 10:5 2-stage array, 4 elements/module (60 elements total)
Post-treatment: Degassing to 5 mg/l dissolved carbon dioxide (dosing to pH 7 with sodium hydroxide)
Spec. energy dem.: 215 kW for 126 m³/h flow, 1.71 kWh m⁻³.
4.3.4 Problem in electrodialysis: energy demand

A light brackish water of the specification below requires demineralisation using a 500 cell pair, \(1 \times 0.75\, \text{m}\) electrodialysis stack for potable water production: \(\text{Na}^+, 830\, \text{ppm}; \text{Cl}^-, 1530\, \text{ppm}; \text{Ca}^{2+}, 100\, \text{ppm}, \text{SO}_4^{2-}, 50\, \text{ppm}; \text{Mg}^{2+}, 60\, \text{ppm}; \text{HCO}_3^-, 120\, \text{ppm}\).

It is desired that the sodium content be halved. If the ED softens at twice the rate of desalination according to first order kinetics, and the process operates with an overall current efficiency of 95%, calculate (a) the total current required, and (b) the specific energy demand in kWh m\(^{-3}\) for a conversion of 85% and an average production rate of 500 m\(^3\)/day desalinated water. How would the specific energy demand change if a further stage were added to yield a 75% desalinated product?

Assume specific cell pair area resistance is given by: \(r\, (\text{ohms cm}^2) = 10 + (10^4/\overline{C})\), \(\overline{C}\) being the average concentration in meq l\(^{-1}\) of the diluate stream passing through the stack.

Solution

Electrodialysis demands a knowledge of the concentration of charge, since this can be directly related to the current demanded by the Faraday equation. Thus all concentrations must be converted to meq/l by dividing by the molar weight and multiplying by the charge:

\[
\begin{array}{c|c|c|c|c}
\text{mg/l} & \text{meq/l} & \text{mg/l} & \text{meq/l} \\
\hline
\text{Na}^+ & 830 & 36 & \text{Cl}^- & 1530 & 43 \\
\text{Ca}^{2+} & 100 & 5 & \text{SO}_4^{2-} & 50 & 1 \\
\text{Mg}^{2+} & 60 & 5 & \text{HCO}_3^- & 120 & 2 \\
\text{Total} & & 46 & & 46 \\
\end{array}
\]

Since one unit of charge will shift a cation and an anion simultaneously, the calculation of the current is based on the total cations or anions removed and not the sum of both.

If hardness (i.e. \(\text{Ca}^{2+}\) and \(\text{Mg}^{2+}\)) is removed at twice the rate of sodium according to first order kinetics, then:

\[
\ln(\frac{C}{C_0})_{\text{hardness}} = 2\ln(\frac{C}{C_0})_{\text{salinity}}
\]

where \(C\) and \(C_0\) are the outlet and inlet concentrations respectively and \(k\) is the first-order rate constant. Thus, the residual/influent hardness is \((\text{residual/influent salinity})^2\).

The composition of the desalinated product in meq/l can then be calculated along with the amount of material removed \((\Delta C)\) and the average concentration of the diluate \((\overline{C})\), respectively from the difference and average of the inlet and desalinated product streams:
From Equation (2.25), the current is given by:

\[ i = \frac{Q F \Delta C}{N \xi} = \frac{500 \times 96 \times 500 \times 25.5}{24 \times 60 \times 60 \times 500 \times 0.95} = 30 \text{ A} \]

From Equation (2.26):

\[ E = i^2 R / \Theta Q \]

where \( R \) is the ED stack resistance in ohms and can be obtained from the cell pair area resistance, the projected membrane area and the number of cell pairs:

\[ R = \frac{rN}{A} = \frac{(10 + 10^4 / 33.2) \times 500}{1 \times 0.75 \times 10^4} = 20.7 \Omega \]

So, the energy demand in kWh per m³ of product at 85% conversion is:

\[ E = \frac{30^2 \times 20.7 \times 24}{0.85 \times 500 \times 1000} = 1.05 \]

Desalination of 75% can be achieved by hydraulic staging (Section 2.4.1), whereby the number of cell pairs is halved and the flow doubled to permit the same current to remove half the quantity of ions as that removed at the first stage. The energy demand for 75% desalination is thus 1.5 times that for 50% desalination at 1.57 kWh m⁻³ for the transfer of ion alone. Pumping of the water, for a pressure drop of 2.5 bar across the stack, adds an additional 0.2 kWh m⁻³, making a total energy demand of 1.77 kWh m⁻³.

It is interesting to compare the relative energy demands of the reverse osmosis (from Section 4.3.1) and electrodialysis processes, which have been calculated on roughly the same basis. For the examples given the specific energy demand values for each process are about the same only if 50% desalination is acceptable, in which case the RO energy demand could be halved by 50:50 blending with feedwater. On the other hand, 85% recovery is readily attainable with the ED, whereas 75% is the maximum possible from a 2-stage RO process. The membrane process choice for light brackish waters such as these may therefore ultimately be made on the basis of both capital cost and specific technical facets of the process (product water purity, operability, footprint, etc.), rather than simply on running costs.

<table>
<thead>
<tr>
<th></th>
<th>( C_0 )</th>
<th>( C )</th>
<th>( \Delta C )</th>
<th>( C_\text{ave} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na⁺</td>
<td>36</td>
<td>18</td>
<td>18</td>
<td>27</td>
</tr>
<tr>
<td>Ca²⁺</td>
<td>5</td>
<td>1.25</td>
<td>3.75</td>
<td>3.13</td>
</tr>
<tr>
<td>Mg²⁺</td>
<td>5</td>
<td>1.25</td>
<td>3.75</td>
<td>3.13</td>
</tr>
<tr>
<td>Totals</td>
<td>46</td>
<td>20.5</td>
<td>25.5</td>
<td>33.2</td>
</tr>
</tbody>
</table>
**4.3.5 Problem in submerged membrane filtration design**

A dead-end hollow fibre submerged microfiltration membrane module having a membrane area $A_m$ of 46 m$^2$ is being used to provide a flow of 600 m$^3$ day$^{-1}$. The process operates at a flux $J$ of 32 l m$^{-2}$ h$^{-1}$ and an aeration rate $Q_a$ of 50 m$^3$ h$^{-1}$ per module, and the water viscosity is around $1.15 \times 10^{-3}$ Pa s.

The resistance of the cleaned membrane is $7 \times 10^{11}$ m$^{-1}$. As a result of fouling by the suspended matter, the operational (reversible) fouling resistance $R_{rev}$ increases by $1.55 \times 10^{11}$ m$^{-1}$ per minute. The residual (irreversible) fouling resistance $R_{irr}$ increases at an average of $5.4 \times 10^9$ m$^{-1}$ per minute over the entire cleaning cycle (i.e. the period between chemical cleans), and the system is chemically cleaned once a pressure of 1 bar is reached.

Backwashing (with permeate) at three times the operating flux for 20 seconds every 12 minutes completely removes the reversible fouling. Cleaning in place by flushing with hypochlorite for 10 minutes and then soaking for a further 40 minutes removes the irreversible fouling.

Calculate (a) the cleaning cycle time, and so (b) the total minimum membrane area requirement and (c) the capital cost, assuming CAPEX, \( £k = £80 + \text{ membrane cost/150} \) given that the membranes cost £40 per m$^2$.

The aeration energy demand (kWh per m$^3$ permeate) is related to the aerator flow by:

\[
E_a = 0.02Q_a/JA_m, \quad \text{for } Q_a > 25 \text{ m}^3\text{ h}^{-1}
\]

under the operating conditions outlined, $Q_a$, $J$ and $A_m$ taking SI units. If electrical costs are 8p per kWh, what is the total electrical energy demand? How significantly is this affected by the backflush energy demand?

**Solution**

The solution proceeds through determination of the pressure changes and operational cycle times from Equation (2.10), and using Equation (2.23) to determine pumping energy demand and then adding the aeration energy to this. The figures must be adjusted to account for downtime and loss of permeate used for backflushing.

The pressure profile is depicted in Fig. 4.23. The filtration backflush cycle pressure change $\Delta P_b$ can be calculated from the rate of increase in the reversible fouling resistance, $R_r = 1.55 \times 10^{11}$ m$^{-1}$ min$^{-1}$, and from Equation (2.10):

\[
P_b = \mu J(R_m + R_r + R_{irr})
\]

where $R_m$, $R_r$, and $R_{irr}$ refer to the hydraulic resistances of the membrane, reversible fouling component and irreversible fouling component respectively. $R_r = R_r t_r$ if the resistance increases linearly with time. Given that $R_{irr} \ll R_r$ over the course of a backflush cycle, the change in pressure over the cycle is:

\[
\Delta P_b = \mu J R_r t_r
\]
So, converting flux to SI units and employing the appropriate viscosity value for 15°C:

$$\Delta P_b = 1.15 \times 10^{-3} \times 6.94 \times 10^{-6} \times 1.55 \times 10^{11} \times 12/10^5 \text{ bar}$$

$$= 0.148 \text{ bar}$$

The cleaning cycle time is also obtained from Equation (2.10). In this case the maximum pressure is given as 1 bar (10⁵ Pa) and $\mu J R_f$ has already been calculated as 0.148 bar. Thus:

$$\mu J (R_m + R_f) = 10^5 - 1480 = 9.85 \times 10^4$$

$$R_f t_f = 9.85 \times 10^5 / \mu J - R_m$$

So,

$$t_f = \frac{1}{5.4 \times 10^9} \left( \frac{9.85 \times 10^4}{1.15 \times 10^{-3} \times 6.94 \times 10^{-6} - 7 \times 10^{11}} \right)$$

$$= 2160 \text{ minutes (36 hours).}$$

Note that if the value for the irreversible fouling rate $R_{ir}$ had applied only to the filtration cycle itself, then $t_f$ would be extended by $100 \times 0.33/12.33 = 2.7\%$ — the ratio of the backflush period $t_b$ (0.33 min) to the total backflush cycle time $t_r + t_b$ (12 + 0.33 min).

The required feed flow rate is dictated by the product flow rate and total downtime. The downtime $t_D$ over the course of a complete cycle is given by the sum of the cleaning period $t_c$ and backflush periods:

$$t_D = t_c + t_b \times t_f / (t_r + t_b) = 50 + 0.33 \times 2160/12.33$$

$$= 108 \text{ min (1.8 h) over the 36 hour cleaning cycle}$$

At a product flow $Q_p$ of 600 m³ per day, this downtime is associated with a volume of:
The total volume demand is further increased by the use of permeate product for backflushing for 20 seconds at three times the flux every 12 minutes which, ignoring the cleaning downtime, equates to a volume demand of:

\[ V_{\text{backf}} = Q_p \times (3 \times 0.33/12.33) \times (1 - t_c/t_{ir}) \]

\[ = 600 \times 0.0811 \times (1 - 50/2160) = 47.5 \text{ m}^3 \text{ day}^{-1} \]

where the \((1 - t_c/t_{ir})\) factor accounts for the cleaning period over which no backflushing takes place.

So, the total volume demand, or feed flow, is:

\[ Q_f = 600 + 30 + 47.5 = 678 \text{ m}^3 \text{ day}^{-1} \]

which implies a conversion of 88%.

For a flux \(J\) of 251 m\(^2\) h\(^{-1}\), the membrane area required is:

\[ A = 678 \times 1000/(25 \times 24) = 1130 \text{ m}^2 \]

Thus the total number of membrane modules required is:

\[ N = A/A_m = 1130/46 = 24.6 \]

which implies 25 membrane modules are required. The capital cost, for membrane costs of £40 per m\(^2\), is then given by:

\[ \text{CAPEX, £k} = £80 + 25 \times 40 \times 46/150 = £386 \text{ K} \]

Operational costs are obtained from the pumping and aeration energy demands. According to Equation (2.23), and based on a 40% pumping efficiency, the liquid pumping energy demand \(E_p\) in kWh per m\(^3\) permeate relates directly to the average TMP (in Pa) over the whole cycle:

\[ E_p (\text{kWh m}^{-3}) = P_{\text{avc}}/(0.4 \times 1000 \times 60 \times 60) = P_{\text{avc, op}}/2.16 \times 10^6 \]

where \(P_{\text{avc}} = (P_{\text{max}} + P_{\text{min}})/2 = (10^5 + \mu R_m)/2 = 52800 \text{ Pa}\).

The pumping demand energy (operating cycle) is thus given by:

\[ E_p = 52800/2.16 \times 10^6 = 0.024 \text{ kWh m}^{-3} \]

At 8p/KWh, this relates to a cost of £0.002 per m\(^3\).

Aeration energy demand is given by:

\[ E_A = 0.02 Q_a/JA = 0.02 \times 50/(25 \times 10^{-3} \times 46) = 0.87 \text{ kWh m}^{-3} \]

which relates to a cost of £0.070 per m\(^3\).
The above costs do not include the energy demanded by the backflush process. Clearly, based on the data given, backflushing cannot impact significantly on the total energy demand of the submerged membrane process since it amounts to only around one-twelfth of the pumping energy demand of the operating cycle, which itself contributes less than 3\% to the total energy demand. Backflushing does, on the other hand, increase the membrane area requirement due to downtime, as has been demonstrated above.

These examples demonstrate the general principles of design. As the dispirate answers from the energy demand calculation by Film Theory (4.3.1) and RO CAD (4.3.3) suggest, basic assumptions made about fundamental properties of the system can profoundly influence the answer obtained. Accurate information on performance, both hydraulic and water purification, demands pilot trials and/or pertinent data from existing full-scale installations. Such data is given in Chapter 5.
Chapter 5

Case studies

Bruce Jefferson
School of Water Sciences, Cranfield University
5.1 Flag Fen high-purity water production plant (UK)

5.1.1 Background

To many the UK is perceived as a water-rich country but the actual situation is geospecific with annual rainfall ranging from up to 1600 mm in the north and west to as low as 550 mm in the south and east. Similar amounts and temporal distributions of rainfall are found in countries such as Israel where water is viewed as a commodity and recycling is commonplace. Since the 1990s legislation has required huge investment in coastal sites to improve effluent quality prior to discharge. Unlike inland water discharges, coastal waters are a lost commodity which offers the potential for reuse without downstream impacts on the water demand balance of the inland areas. Specifically, with further treatment the coastal discharges could prove an excellent source of industrial water intake. Not only does this provide a reliable source of water to the industrial client but alleviates a commensurate amount of potable water stress on the public supply.

Anglian Water Group (AWG) is the water and sewerage operator for a region of the UK in the east of the country and suffers from some of the lowest annual rainfall in the country. Increasing population growth within the AWG supply region has led the company to look at the strategic benefits of the use of coastal discharges. A survey of potential industrial partners was conducted based on the following criteria:

- Proximity to a sewage works
- Proximity to a coastal discharge
- Non-food industry
- Need for significant amounts of ultrapure water
- Long asset life (> 10 years)

One of the ideal candidates was identified as the power station in Peterborough owned by Texas Utilities (TXU). The station is located adjacent to Flag Fen sewage works which treats the majority of the flow from Peterborough. Negotiations between TXU and Alpheus Environmental (a wholly owned subsidiary of AWG) determined the contractual and expected product water quality from the advanced treatment works that would supply the power station (Table 5.1). The plant was built in the spring of 2000 and started operating in the summer.

5.1.2 Description of plant

The initial process selection was controlled by the requirement to produce high purity water with a conductivity of 70 $\mu$S cm$^{-1}$ which restricted the final stage to a reverse osmosis plant. The real challenge of the project concerned the pretreatment to the RO plant which had to provide a robust feed water from a
sewage effluent which has a naturally variable effluent quality. A similar situation exists in the USA at Water Factory 21 where a multiple treatment train is used. This comprises of pH adjustment to 10, coagulation, super chlorination, clarification, depth filtration, pH adjustment to 6 and finally dechlorination. This approach suffers from a number of drawbacks including a large chemical use, large footprint and high operating costs. The original process train also included activated carbon but was discontinued due to fines coming out of the filter beds and blocking the RO tubes. As an alternative to the above, AWG investigated the use of microfiltration membranes. Previous research had shown membrane processes to be a suitable unit operation for the removal of solids, bacteria and microorganisms capable of robustly producing a silt density index below 2. Four commercial systems were tested including the specific pre-screening requirements which ranged from 1 mm screens to sand filters (~50 μm equivalent screen).

The final process train selected consisted of a pre-screen, microfiltration plant and a reverse osmosis plant (Fig. 5.1). Wastewater from the screens and the microfiltration plant is returned to the head of the sewage works. Concentrate disposal is achieved by dilution with final effluent from the sewage works in the final effluent chamber and discharged into the tidal section of the river Nene.

Pre-screens

Appropriate pre-screening was found to be important to ensure suitable operability of the plant. Assessment of the different options resulted in a Hydrotech HDF 830 drum screen being installed. The screen removes solids larger than 150 μm and has a self-cleaning facility to remove the retained solids by back flushing with screen effluent. The screen is manufactured from stainless steel with a polyester cloth insert material. To ensure constant flow to the membrane plant the screens were fitted with variable speed pumps which controlled the level in the screened water tank.

Microfiltration

The screened effluent then passes to the microfiltration plant. The plant comprises two parallel streams, each containing 17 modules. Each module contains 50 m² of PVDF hollow fibre membrane operating in a semi dead-end mode with the flow from outside to in. The membrane material is rated with a nominal pore size of 0.1 μm and fabricated into individual membrane units with

### Table 5.1 Contractual water quality

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>Contractual</th>
<th>Expected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conductivity</td>
<td>μS cm⁻¹</td>
<td>70</td>
<td>50</td>
</tr>
<tr>
<td>Silica dissolved</td>
<td>mg ¹⁻SiO₂</td>
<td>1</td>
<td>0.5</td>
</tr>
<tr>
<td>Sodium</td>
<td>mg ¹⁻Na</td>
<td>20</td>
<td>15</td>
</tr>
<tr>
<td>Calcium</td>
<td>mg ¹⁻CaCO₃</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>TOC</td>
<td>mg ¹⁻C</td>
<td>2.5</td>
<td>&lt;2</td>
</tr>
<tr>
<td>Polysaccharides</td>
<td>µg ¹⁻</td>
<td>100</td>
<td>50</td>
</tr>
</tbody>
</table>
a height of 2000 mm and an outside diameter of 150 mm. The plant has a total membrane area of 1700 m² and can produce up to 1500 m³ d⁻¹ (Table 5.20). The membranes are operated at an average flux of 30.5 LMH and a transmembrane pressure range between 0.4 and 3 bar yielding an overall recovery of 90%. Membrane fouling is controlled through a sequence of cleaning cycles. An automatic air/water backwash is conducted every 12 minutes for 90 seconds. In addition each stream is chemically cleaned with a hypochlorite and pH adjustment for 90 minutes every 14 days through an automatic cleaning in place (CIP) system. Membrane integrity is monitored by particle counters and turbidity with additional pressure decay tests conducted periodically.

Reverse osmosis
The final stage of the treatment train is two parallel reverse osmosis units arranged in a 4:2 array. The membranes are polyamide spiral-wound modules supplied by Koch Membrane Systems (UK) and are rated at a molecule weight cut off of 200 kDa. The plant contains a total membrane area of 1451 m² and is designed to produce up to 1200 m³ d⁻¹ of high-purity water at a recovery of approximately 80%. The membranes operate at a pressure range between 6 and 16 bar producing an average flux of 34.5 LMH. Membrane cleaning is triggered by a 15% reduction in the monitored variables and when necessary initiates a 360-minute clean with high and low pH solutions to remove both organic and inorganic fouling. Membrane integrity is monitored by checking conductivity of the membrane banks and individual pressure vessels.

5.1.3 Performance
The major challenge with using a sewage effluent as the feed to a high-purity treatment plant is the natural variation in its quality which varies both seasonally and throughout the day. Flag Fen produces an effluent water quality with a mean BOD₅ of 10 mg l⁻¹, COD of 51.3 mg l⁻¹ and a total suspended solids concentration of 17 mg l⁻¹. The quality is variable with a peak to mean ratio of
3.7 for suspended solids and 3.3 for BOD₃. Smaller variations are observed for COD (2.2) and total phosphorus (1.5). Seasonal variations occur due to a vegetable processing factory that discharges effluent of very different quality depending on the food being processed.

The permeate quality from both membrane plants was both reliable and in line with previous values found from pilot plant work. Feed quality to the RO plant can show large variation with rainfall; for instance, a conductivity of between 800 and 1200 μS cm⁻¹. Permeate quality from the RO plant is 38.5 μS cm⁻¹ representing a rejection of 96.7% on average. Overall rejection of TDS is 93% with specific rejections of sodium and calcium of 95 and 99% respectively (Table 5.20).

The main issue of concern was membrane fouling reducing output from the MF plant and increasing pressure on the feed stream of the RO plant. Initial operation of the plant showed excellent water quality but high RO fouling rates. The RO feed pressure increased at a rate of 1 bar per hour at worst due to rapid build up of calcium phosphate. Changes to upstream operation has effectively controlled the problem and membrane cleaning is now required every 6 months on average.

The Flag Fen plant is currently operating well and exceeding the target water qualities set down in the original negotiations. The quality of the delivered water has enabled ultrapure water production at the power station to increase by 20%. Coupled to this is a reduction of over 90% in the costs of ion exchange regeneration due to an increase in the operating cycle of the twin bed demineralisation plant from 8 to 60 hours. Overall, 1250 m³ of tap water per day have been saved which has reduced the station’s total water use by 11%. Client receptivity to the scheme is very high as supply is guaranteed and operating costs decreased. The high-purity water plant cost around £1 000 000 ($1 524 000) and each membrane stage has an operating cost of 7.5p m⁻³ ($0.11 m⁻³). The success of the scheme has attracted media attention and the project has won a number of prestigious awards including the 2000 Water UK/Environment Agency water, the 2001 IChemE innovation efficiency award and the 2001 Green apple award.
5.2 Eraring Power Station: purification of secondary sewage for boiler feedwater (Australia)

5.2.1 Background

Pacific Power's Eraring plant is a 4 $\times$ 660 MW coal-fired power station, situated on Lake Macquarie which provides about 25% of the power requirement to New South Wales. The power station utilises salt water from the lake for condenser cooling with all other water supplied as potable by Hunter Water Corporation. Prior to 1988 potable water usage was almost 8.5 Ml d$^{-1}$ and represented one of the three largest costs to the power station. To reduce costs a water audit was conducted to identify the major areas of use. After engineering modifications the water use was reduced to 4 Ml d$^{-1}$:

- In the chemical control room, where various water samples are automatically analysed, the supply was modified from using potable water to using recirculating auxiliary cooling water.
- In the ash and dust system, gland sealing water was changed over from potable water to salt water. This increased costs in the ash and dust area but the saving in water costs overshadowed the increased maintenance.
- The wash down systems were modified to small high pressure nozzles.

The audit indicated that significant savings could be made if another supply of process water could be found. A source became available when the Hunter Sewage Project upgrade of the local Dora Creek sewage treatment plant was initiated. As part of the upgrade a new wastewater treatment plant was to be built at Dora Creek which is approximately 6 km south of the power station. The sewage effluent was to be pumped under Lake Macquarie to an extended ocean outfall. The discharge pipeline was due to pass close to the power station and so reuse became an option.

Following a detailed review period, a deed of agreement was signed between Pacific Power and Hunter Water Corporation for a guaranteed 15-year period. During this period Pacific Power must accept all secondary effluent up to a maximum of 5.2 Ml d$^{-1}$. The effluent quality from the Dora Creek sewage works (Table 5.2) means that additional treatment is required prior to use at the power plant. The principal use of processed water is for demineraliser feed with the remaining flow being used for other water applications on site:

- Demineralising plant (1.5 Ml d$^{-1}$)
- Auxiliary cooling towers (1 Ml d$^{-1}$)
- Wash down water
- Fire services
- Ash disposal system (1–1.5 Ml d$^{-1}$)
### Table 5.2 Specifications for water quality

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Maximum value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suspended solids (mg l⁻¹)</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Silt density index</td>
<td>&lt; 3</td>
</tr>
<tr>
<td>Reovirus (100 ml⁻¹)</td>
<td>Nil</td>
</tr>
<tr>
<td>Enterovirus (100 ml⁻¹)</td>
<td>Nil</td>
</tr>
<tr>
<td>Total coliforms (100 ml⁻¹)</td>
<td>&lt; 10</td>
</tr>
<tr>
<td>Faecal coliforms (100 ml⁻¹)</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Faecal streptococci (100 ml⁻¹)</td>
<td>&lt; 1</td>
</tr>
</tbody>
</table>

Demineralising feed water has two requirements. The first is that the water must not contain any organic matter that will foul the ion exchange resin, particularly anion resin. The second is that the TDS should not impose any additional load on the plant over that of domestic feedwater. Benefits can be easily obtained if the TDS is in fact lower than the original supply as it can significantly reduce costs through fewer regenerations of the demineraliser plant. A prime requirement for all end uses, and one required by the New South Wales Environmental Protection Agency, is that the water be disinfected (Table 5.2). A number of technologies were considered including ponding, wetlands and UV disinfection but all required additional treatment before the water was suitable for reuse at the plant and so membranes were ultimately used.

#### 5.2.2 Description of plant

Secondary effluent is supplied from the Dora Creek sewage treatment plant and is initially stored in a 8 Ml holding tank before being transferred to the treatment plant. The flow is supplied at a rate of 3.5 Ml d⁻¹ and blended with tertiary effluent from the station’s sewage works and contaminated plant water which has had the oil and grit removed. The flow then passes through a motorised screen before being pumped to the microfiltration plant (Fig. 5.3).

Filtrate from the MF plant is dosed with sodium hypochlorite en route to a storage tank to control biological growth. Sulphuric acid (4%) is also added to reduce pH and minimise hydrolysis of the RO membrane. Water is then pumped from the storage tank, dosed with anti-scalant, and screened through a 5 μm disposable cartridge filter before passing to the RO plant. Permeate is pumped to a degasser unit before being fed preferentially to the demineraliser plant because of its low TDS. The RO reject stream is dosed with ferrous chloride before being passed to the station’s ash dam together with waste from the demineraliser plant and the wastewater sump.

**Microfiltration**

The microfiltration plant incorporates two parallel streams each containing 90 modules. Each module houses 15 m² of polypropylene hollow fibre membrane supplied by Memcor and rated at a nominal pore size of 0.2 μm. The plant
contains a total membrane area of 2700 m² and is designed to treat 5.2 Ml d⁻¹ at an overall recovery of 90%. The membranes are supplied by pumps operated at a pressure of 450 kPa (4.5 bar) which delivers an average flux of 72 LMH (calculated from data above). Membrane fouling is controlled through a sequence of cleaning cycles and is triggered on either TMP or differential pressure. The membranes are initially drained and then high pressure air (600 kPa (6 bar)) is blown through the membranes to loosen attached material followed by a back pulse of permeate. The backflush cycle occurs between every 17 and 60 minutes depending on fouling. In addition, every 200 service hours a CIP is carried out with caustic soda (1%) and a detergent. The cleaning solution is reused on average for 10 cycles before being replaced. Membrane integrity of an array is monitored through an automatic pressure decay test which is carried out every 24 service hours. When necessary the individual module can then be identified through a sonic test (out of service). Air is supplied at 100 kPa (1 bar) and if noise is detected the individual module is isolated from the array. When necessary to ensure production the modules are pin repaired where any broken fibres are plugged using a stainless steel pin at both ends of the module.

The final stage of the treatment train is two parallel reverse osmosis units arranged initially in a 6:3 array (later converted to a 10:4 array). The membranes are cellulose acetate spiral wound modules and are rated at a 98% salt rejection. The plant is designed to produce up to 3.75 Ml d⁻¹ of high-purity water at a recovery of approximately 80%. The membranes operate at a pressure range between 1500 and 3500 kPa (15–35 bar) producing an average flux of 13 gfd (22 LMH). The plant is monitored in terms of the normalised permeate flow (temperature, pressure and concentration) and under normal operation is measured at 13–151 s⁻¹. Chemical cleaning is triggered by a reduction of 15% in the normalised flow rate.
5.2.3 Performance

The plant is reported to have performed well since its installation and has resulted in significant water savings. The percentage of reclaimed water used has increased gradually from 5.2% during installation in 1994 to 56.7% in 1998 such that up to February 1999 the plant has used a total of 2,333 Ml of reclaimed water. In 1999, potable water use at the site was 1.8 Ml d\(^{-1}\) and this is expected to be ultimately reduced to 400 kl d\(^{-1}\) by 2010 when the duration of the deed of agreement is complete.

The treatment performance is indicated by a reduction in BOD\(_5\) from 20 to 50 to less than 1 mg l\(^{-1}\) across the whole plant (Table 5.20). The equivalent removal for turbidity is from 50 to <0.1 NTU and for Faecal coliforms from <10\(^6\) to <1. RO permeate is also low in dissolved solids with a permeate concentration of specific ions of 32 mg l\(^{-1}\) (Cl), 2.2 mg l\(^{-1}\) (Si), 17.8 mg l\(^{-1}\) (Na) and 0.5 mg l\(^{-1}\) (Ca). The low levels of dissolved solids have increased demineraliser cation operation capacity from 2176 m\(^3\) with potable water to 4792 m\(^3\) with reuse water and anion operation capacity from 2113 m\(^3\) to 3472 m\(^3\) respectively before requiring a regeneration.

The main operational concerns have been periods of increased membrane fouling. At one stage the MF plant was not responding to the cleaning cycle. Diagnosis revealed manganese fouling was occurring which was completely ameliorated with a citric acid (citriclean) clean. Similarly, at one stage RO cleaning frequency increased dramatically. The problem was linked to organic fouling which decreased naturally as indicated by reduced chlorine demand and ammonia levels in the plant. If the problem occurs again a chlorine chemical clean is planned to oxidise the organic layer. An important aspect of the scheme has been gaining employee acceptance of using reclaimed water. To allay fears and concerns regular testing for bacteria and viruses is reported and personal protective equipment (ppe) and covers for equipment is supplied were appropriate.

The water reclamation plant originally required a total capital cost of AUD\$4.5 million ($3.34 million) in 1994 of which AUD\$4 million ($2.96 million) was construction and commissioning and AUD\$0.5 million ($0.37 million) was required for segregating the potable and reclaimed water supplies. A further
AUD$180 000 (S133 560) was required to upgrade the throughput from 2.5 ML d\(^{-1}\) to 3.75 ML d\(^{-1}\) in December 1998.

The operating costs of the schemes consists of chemicals, spare parts, analytical costs, effluent supply and the service agreement payments (Table 5.3). The chemical costs are generated from cleaning the MF plant (17.5%), RO pretreatment (58%), RO cleaning (1%) and reject treatment (14%). The overall opex has remained relatively stable during the initial 5 years of operation where increasing spares costs have been balanced by reduced supply and contract charges as the plant output increases. The service contract is negotiated at the end of 1999 when it is expected to increase and so impact on the overall opex of the plant.

The reclamation plant generates two major savings for Pacific Power. Firstly saving based on reduced potable water use on site which increased from AUD$78 500 ($58 247) in 1994/95 to AUD$726 200 ($433 904) in 1998/99 (Table 5.4). The savings are expected to continue to increase once the plant is at full capacity, generating an annual saving of AUD$1 100 000 ($588 720). An

<table>
<thead>
<tr>
<th>Table 5.3</th>
<th>Production costs (AUD$ m(^{-3}))</th>
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<tbody>
<tr>
<td>Analytical</td>
<td>0</td>
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<tr>
<td>Spares</td>
<td>0</td>
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<tr>
<td>Contract</td>
<td>0.107</td>
</tr>
<tr>
<td>Chemicals</td>
<td>0.208</td>
</tr>
<tr>
<td>Supply</td>
<td>0.017</td>
</tr>
<tr>
<td>Total</td>
<td>0.332</td>
</tr>
<tr>
<td>Total ($ m(^{-3}))</td>
<td>0.246</td>
</tr>
</tbody>
</table>

Currency conversion (x 0.5975–0.778) based on 12 August 1994–1998 figures (http://www.oanda.com/convert/fxhistory (accessed November 2002)).

<table>
<thead>
<tr>
<th>Table 5.4</th>
<th>Cash balance of reclamation plant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reclaimed water (ML)</td>
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</tr>
<tr>
<td>Saving (AUD$)</td>
<td>78 500</td>
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<tr>
<td>Demin. regeneration (no.)</td>
<td>51</td>
</tr>
<tr>
<td>Saving (AUD$)</td>
<td>26 800</td>
</tr>
<tr>
<td>Total saving (AUD$)</td>
<td>105 300</td>
</tr>
<tr>
<td>Total saving ($)</td>
<td>78 132</td>
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<tr>
<td>Total cost (AUD$)</td>
<td>44 238</td>
</tr>
<tr>
<td>Net saving (AUD$)</td>
<td>61 062</td>
</tr>
<tr>
<td>Net saving ($)</td>
<td>45 308</td>
</tr>
</tbody>
</table>

Currency conversion (x 0.5975–0.778) based on 12 August 1994–1998 figures (http://www.oanda.com/convert/fxhistory (accessed November 2002)). Total costs based on reclaimed water amount and quoted cost per m\(^3\) and recoveries of 80% and 90% across the MF and RO respectively.
additional benefit is a reduction in operating cost of the demineralising plant through a reduced number of regenerations. The saving predicted when the plant is at full capacity is AUD$100 000 ($53 520) generating a total annual saving of AUD$1 200 000 ($642 240).

The fixed cost of supply from the deed of agreement is a crucial aspect of the economics as it generates significant cost reductions over the period of the deed. The expected payback period on current numbers is between 6 and 7 years enabling 8–9 years of annual saving of around AUD$120 000 ($642 240) as profit. The total saving over the period of the deed equates to AUD$11 100 000 ($5 940 720). The deed agreement was possible as Hunter Water Corporation saved AUD$2 700 000 ($1 445 040) in expenditure on disposal pipeline.

5.3 Doswell combined cycle power plant: zero liquid discharge (USA)

5.3.1 Background

The Doswell combined cycle facility is a 660 MW power plant owned by Doswell Limited Partnership and operated by Bechtel and is one of the largest independent power plants in the USA. The plant is designed for dispatchable load operation working mainly on weekdays during the winter and summer and contains two parallel units which share a common water treatment plant. The plant is designed to burn natural gas and has been in operation since July 1991.

A major concern about the plant was potential limitations on water supply and wastewater discharges issued by the local government of Hanover, Virginia. To address these issues the company decided to minimise water use and recycle wastewater. As part of this process a number of conservation and reuse measures were implemented at the site:

- Air-cooled condensers were installed instead of the typical wet cooling towers to eliminate water losses through evaporation and dramatically reduce wastewater production.
- Dry hybrid burners were installed that limited NOx formation without the need for water or steam injection.
- Potable water demand was reduced to a minimum by utilising sewage effluent from the local wastewater treatment facility.
- Any wastewater generated at the site was recycled through a zero liquid discharge (ZLD) facility.

A key unit process of ZLD is vapour compression evaporation which can recover about 95% of the waste stream as distillate. The remaining flow is then converted into solids in a crystalliser/dewatering device. However, the process is very expensive when flows are high and/or low in dissolved solids. Consequently, the company investigated the possibility of pre-concentrating the flow with...
membrane technologies. Ultimately, a combination of electrodialysis reversal (EDR) and reverse osmosis was adopted to pretreat the waste flows prior to evaporation.

5.3.2 Description of system

The water treatment plant is made up of three integrated treatment systems designed to meet boiler feed water/steam purity and a ZLD requirement (Fig. 5.5):

- Raw water pretreatment
- Boiler feedwater (make up water) treatment
- Wastewater treatment

The plant takes water from both the wastewater and potable water facilities at Hanover county and discharges only solids in the form of filter cakes from both the pretreatment and wastewater treatment plants.

Raw water pretreatment

The raw water pretreatment plant is designed principally for solids removal from the incoming Hanover county sewage effluent (grey water), backwash water and wastewater from the oily water collection system. Raw water enters a coagulation/flocculation chamber followed by a clarifier and dual media depth filters. Backwash water from the filters is periodically returned to the clarifier. Clarifier sludge is dosed with polymer before being thickened and then sent to the filter press for dewatering. The cake is sent to landfill and the recovered water returned to the clarifier.

Make up water treatment

Treated raw water is mixed with potable water and pumped to the boiler feedwater treatment system. The system is designed to remove 99% of the

Figure 5.5 ZLD strategy at Doswell combined cycle power plant
dissolved minerals and provide high-purity water to the boiler. The mixed water flows through a reverse osmosis plant operating at a recovery of 80% and an average salt rejection of 95%. Permeate from the RO mixes with product water from both the waste RO unit and the distillate from the brine evaporator/crystalliser situated in the wastewater treatment plant. The combined flow then enters a degasifier, to remove carbon dioxide, and a mixed bed dimineraliser. The mixed bed plant consists of two 100% capacity ion exchange vessels which remove the final 5% of the dissolved salts. The ion exchange beds process 2 200 000 gallons (8 327 m³) before being regenerated. Waste from the process is pH adjusted and combined with the RO reject before being pumped to the wastewater treatment plant.

Wastewater treatment
The wastewater treatment plant is designed to treat 250 gpm (56.8 m³ h⁻¹) of which 66% is recovered by the membrane processes and the rest through the brine evaporator/crystalliser unit (Fig. 5.6). The wastewater flow is generated by make-up RO reject (64%) (from make-up water plant), power block blowdown (22%) and mixed bed regenerate waste (14%). The combined wastewater flow initially passes through two 100% flow dual media anthracite/sand depth filters operating in a duty standby/backwash mode. Filter permeate is then treated in an EDR unit containing micron feed filters and three 50% capacity membrane stacks. Each parallel line contains 3 stacks in series consisting of 500 pairs of cation- and anion-selective membranes. The EDR unit is designed to recover 84% of the flow with the remaining 16% being sent to the brine tank. The EDR unit includes acid injection for pH control, anti-scalant and clean in place systems to control fouling. The three stages in each stack are operated at voltages of 299, 344, and 264 V with corresponding currents of 17, 11 and 4.8 amps.
respectively. The feed pump discharges at a pressure of 81 psi (5.6 bar) with a differential pressure across the stacks of 14 psi (0.96 bar) on the positive side and 18 psi (1.24 bar) on the negative.

The flow then enters a reverse osmosis plant containing three parallel streams designed at 50% flow enabling continuous operation. Each stream contains 24 cellulose acetate membranes arranged in a 4:2 array. The plant operates at an overall recovery of 75% and a salt rejection of 95%. Permeate is pumped to the demineralisation storage tank and reject is sent to the brine storage tank where it is mixed with the EDR reject.

Treatment of the brine is conducted in a vertical tube, falling film evaporator driven by vapour compression. Wastewater is pH adjusted to between 5.5 and 6 and then heated to boiling point and deaerated. Hot brine then enters the evaporator sump where it mixes with recirculating brine slurry which is pumped to the top of 2 inch (50.8 mm) heat transfer tubes. As the slurry falls a small portion of the water evaporates and condenses on the outside of the heat transfer tubes. The brine evaporator recovers 95% of the flow which is passed on to the demineralisation feed tank with a water quality of less than 10 ppm TDS. The 5% concentrated brine then enters a crystalliser where a further 95% of the remaining water is recovered. The stream is finally sent to a filter press and dewatered to a 20% moisture content sludge which is disposed off site.

5.3.3 Performance

Inclusion of the EDR/RO pretreatment stage reduces the design flow of the brine evaporator from 247 gpm (56 m³ h⁻¹) to 89 gpm (20.2 m³ h⁻¹). The EDR unit

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*Figure 5.7  Ionics Aquamite XX EDR unit for Doswell combined cycle facility*
effectively reduces the concentration of all dissolved ions in the flow (Table 5.5). Total dissolved solids are reduced from 1612 mg l\(^{-1}\) to 217 mg l\(^{-1}\), generating a concentrated stream of 10910 mg l\(^{-1}\). Specific ionic removals range from 47% (fluoride) to 96% (magnesium).

Evaporation is a key unit process in achieving ZLDs due to its ability to operate at high recovery rates from very high TDS waste streams. The technology is however very expensive and as such an economic driver exists for pre-concentrating the flow in systems like the EDR/RO described above. The EDR/RO system cost $750 000 to install and resulted in a 64% reduction in the required capacity of the brine evaporator. The reduction in required capacity of brine treatment results in a saving of $900 000 in capital and $682 day\(^{-1}\) in operational costs (including EDR/RO costs) ($240 000 year\(^{-1}\)). The EDR/RO costs (Table 5.6) include replacement membranes for the EDR in 10 years and the RO in 2 years. The operating costs of the process equate to $0.12 per 1000 gallon ($0.03 m\(^{-3}\)) for the EDR and $0.1 per 1000 gallons ($0.026 m\(^{-3}\)) for the RO. Additional costs savings are made from recycling clean water which reduces costs and make up water demand.

### 5.4 VHP Ugchelen: paper mill water recycling (Netherlands)

#### 5.4.1 Background

VHP security paper mill owned by Ugchelen BV, located in Apeldoorn (Netherlands), produces bank notes and other security papers. The paper mill uses cotton as its raw material which it bleaches with hydrogen peroxide at a temperature of approximately 100\(^{\circ}\)C and pH values between 11 and 12. The process uses a total of 100 m\(^{3}\) of water per ton of paper which at the plant’s

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Feed</th>
<th>Product</th>
<th>Brine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium (mg l(^{-1}))</td>
<td>23</td>
<td>1.2</td>
<td>159</td>
</tr>
<tr>
<td>Magnesium (mg l(^{-1}))</td>
<td>4</td>
<td>0.17</td>
<td>25</td>
</tr>
<tr>
<td>Sodium (mg l(^{-1}))</td>
<td>504</td>
<td>69</td>
<td>3510</td>
</tr>
<tr>
<td>Potassium (mg l(^{-1}))</td>
<td>21</td>
<td>2</td>
<td>160</td>
</tr>
<tr>
<td>Barium (mg l(^{-1}))</td>
<td>0.021</td>
<td>&lt;0.002</td>
<td>0.16</td>
</tr>
<tr>
<td>Bicarbonate (mg l(^{-1}))</td>
<td>80</td>
<td>38</td>
<td>207</td>
</tr>
<tr>
<td>Sulphate (mg l(^{-1}))</td>
<td>898</td>
<td>57</td>
<td>6248</td>
</tr>
<tr>
<td>Chloride (mg l(^{-1}))</td>
<td>153</td>
<td>38</td>
<td>821</td>
</tr>
<tr>
<td>Fluoride (mg l(^{-1}))</td>
<td>1.13</td>
<td>0.59</td>
<td>4.1</td>
</tr>
<tr>
<td>Nitrate (mg l(^{-1}))</td>
<td>82</td>
<td>12.8</td>
<td>137</td>
</tr>
<tr>
<td>TDS (mg l(^{-1}))</td>
<td>1612</td>
<td>217</td>
<td>10910</td>
</tr>
<tr>
<td>Conductivity</td>
<td>2398</td>
<td>333</td>
<td>13592</td>
</tr>
<tr>
<td>pH</td>
<td>6.7</td>
<td>6.5</td>
<td>7.2</td>
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<tr>
<td>Total hardness (mg l(^{-1}))</td>
<td>75</td>
<td>3.5</td>
<td>490</td>
</tr>
<tr>
<td>TOC (mg l(^{-1}))</td>
<td>6.2</td>
<td>4.3</td>
<td>22.6</td>
</tr>
</tbody>
</table>
Table 5.6  Cost balance for different brine treatment options (based on 1990 data)

<table>
<thead>
<tr>
<th>Cost</th>
<th>250 gpm evaporator</th>
<th>90 gpm evaporator</th>
<th>Crystalliser</th>
<th>EDR/RO system</th>
</tr>
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<tbody>
<tr>
<td>Capital and installation ($)</td>
<td>3 600 000</td>
<td>1 950 000</td>
<td>1 370 000</td>
<td>750 000</td>
</tr>
<tr>
<td>Energy ($ day⁻¹)</td>
<td>1471</td>
<td>535</td>
<td>148</td>
<td>300</td>
</tr>
<tr>
<td>Maintenance ($ day⁻¹)</td>
<td>100</td>
<td>54</td>
<td>38</td>
<td></td>
</tr>
</tbody>
</table>

production rate of 5000 tons per year equates to a water usage of 500 000 m³ year⁻¹. The production process results in a high strength wastewater (COD = 4390 mg l⁻¹) at a temperature of 75–85°C and a high pH. Traditionally the effluent from this process has been discharged into the sewer and the water supplied from a local spring requiring 1.5 × 10⁶ m³ of gas to heat it to the required temperature annually.

European legislation is increasing regularly such that paper mills are being forced to minimise their energy and water consumption. The driver for reuse in this case was the combination of compliance with current and near-future legislation together with the financial benefits of reduced water intake, discharge costs and gas supply. One of the most important issues in this specific case was the reuse of both the water and heat from the process.

5.4.2 Description of system

Flow from the production facility initially enters a storage tank prior to passing on to a dissolved air flotation plant at a rate of 10 m³ h⁻¹ (Fig. 5.9). Bubble production is achieved by releasing a super saturated solution of water and carbon dioxide-rich air. This has the added benefit of controlling the pH such that at a ratio of 7 m³ of gas to every 1 m³ of wastewater the pH decreases from 11 to 8.2.

The flow then enters a side-stream membrane bioreactor with a working volume of 200 m³ and a hydraulic retention time of 22 hours. The bioreactor is operated at a MLSS of approximately 20 000 mg l⁻¹, equating to a sludge production of less than 0.02 kgSS d⁻¹. The bioreactor is configured with a side stream loop of 8 mm tubular PVDF membranes rated at a pore size of 0.04 μm. The MBR is designed to treat an average flow of 9 m³ h⁻¹ and contains 82.5 m² of membrane configured in 3 trains of 6 modules. The membranes are operated at a mean TMP of 3.5 bar delivering a mean flux of 120 LMH at a cross-flow velocity of 3.5 m s⁻¹. The membranes have not required cleaning in the first year of operation but when necessary they will be cleaned by a mixture of anionic and cationic surfactants. The membranes have an expected life of 3 years but with the minimal cleaning required so far it is hope this may be significantly extended.

The plant includes an internal heat exchange loop whereby hot incoming flow is cooled to the required temperature for bio treatment whilst at the same time increasing the temperature of the product water stream. This has a direct benefit of reducing the heat required to produce water at the required temperature for processing the raw cotton.
The initial phase of the work included a pilot-scale investigation of thermophilic MBRs for treatment of the wastewater. The performance of the bioreactor was seen to be temperature-dependent with an optimum performance of over 85% removal of COD occurring at 50°C. This equates to an effluent concentration of 600 mg l\(^{-1}\). The effluent was then tested for its suitability for reuse at the plant and was seen to increase the bleaching requirement by 10% compared to the use of groundwater.

The full-scale plant went into operation in 2001 and effluent results for the first 100 days of operation have indicated that even better performance is possible with an effluent COD ranging between 450 and 500 mg l\(^{-1}\) with the reactor temperature at 60°C. Furthermore, COD values are still decreasing indicating that the biomass could still be adapting to the specific conditions of the reactor. The effluent is currently used for 80–90% of the bleaching process without any noticeable deterioration in the product quality.

The utilisation of a reuse system at the site has had significant impact of the flows of fresh and wastewater. The freshwater intake for bleaching has decreased by 80% from 10 to 2 m\(^3\) ton\(^{-1}\) which equates to an annual saving in freshwater
of 40,000 m³. The internal heating loop has reduced gas consumption by 20% from 660 m³ ton⁻¹ to 520 m³ ton⁻¹ producing an overall saving in gas of 700,000 m³. Total wastewater discharged from the sites has halved to a population equivalent of 5000 p.e. which generates a saving of €227,000 per year ($226,000).

5.5 Kronospan Ltd: medium-density fibreboard wash water recycling (UK)

5.5.1 Background

The production of medium-density fibreboard (MDF) involves a number of simple but large-scale operations. Forest thinning and sawmill residues are debarked and chipped before being washed to remove residual dirt and grit from the wood. The fibres are then steam-softened, cooked and refined between two flat plates and finally mixed with resin to produce the fibreboard.

An example facility is at Chirk in North Wales where Kronospan UK manufacture chipboard, MDF, melamine-faced boards and sawn timber. The plant produces 47,800 m³ of MDF effluent per year of which the majority is associated with the washing and refining stages of production. The quantity of excess water generated during the process depends on the moisture content of the wood and can range from 400 l per tonne of bone dry wood processed in the summer to up to 600 l in the winter. The characteristics of the effluent are high COD and high suspended solids of which cellulose, lignins and resin acids are key components.

Prior to 1995 the effluent was tankered off site at a cost of £9.8 tonne⁻¹ ($14.72 tonne⁻¹). The company was not only concerned about the cost of this option but also the risk to production if the tankers failed to arrive as the site could only store one day’s effluent. A decision was made to incorporate on-site treatment at the production facility which should meet the following aims:

- Low capital cost/rapid investment payback
- Optimum product/resource recovery
- Effluent reduction, recycling and reuse
- Long-term environmental compliance
- Easy to upgrade modular system

The site contained an existing conventional biological treatment train comprising a dissolved air flotation unit followed by an activated sludge plant. Problems with effluent reliability and limited resource recovery due to the poor effluent quality led to the search for an alternative solution. Ultimately, a treatment train containing membranes was installed in June 1995, which offered the potential for zero discharge of the wash water effluent.
5.5.2 Description of plant

The effluent treatment plant treats all the MDF effluent generated within the production facility. Excess flow is stored prior to it being pumped into the treatment plant. The plant consists of multiple treatment stages combining physicochemical processes with membrane technology (Fig. 5.10). Water entering the plant is initially dosed with a polyelectrolyte and flocculated. The flocculated suspension is then pumped into a plate and frame filter press from which a 45–50% dry solids cake is recovered. The filter cake can be burnt in the boiler or re-used in chipboard production. The filtrate passes through a dual media depth filter before entering a four-stage reverse osmosis plant. Permeate from the membrane plant is polished through an activated carbon bed before being reused on site for boiler feed. Concentrate from the membrane plant is returned to the resin make up tank and backflush/wash water from the sand filter and filter press are combined and returned to the head of the treatment works.

Pretreatment

Polyelectrolyte is dosed up to a rate of 500 mg l⁻¹ into the flow and then flocculated in a 15 m³ tank with a residence time of 30 min. The project included the development of a new polyelectrolyte specifically designed for the needs of MDF effluent. The flocculated suspension then passes to a 55-plate filter press at up to 50 m³ h⁻¹ and 6 bar (treating an average flow of 30 m³ h⁻¹). The press (Fig. 5.11a) produces up to 2 m³ of 45–50% solids cake per batch. Filtrate from filter press then passes to a dual media depth filter containing sand and anthracite. The filter operates at a design velocity of 6 m³ m⁻² h⁻¹ and requires cleaning with a combined air scour and water backwash which results in a water

Figure 5.10  Process flow diagram for Kronospan zero discharge and recovery plant
recovery across the bed of 98–99%. Finally the flow passes through a 5 μm cartridge filter to remove any gross solids and filter grains that might pass through the bed.

Reverse osmosis
The centre of the treatment plant is a single stream reverse osmosis plant configured in a four-stage feed and bleed array, each with its own cross-flow recirculation pump. The membranes are polyamide 8040 spiral-wound modules developed with Osmonics. The plant contains a total membrane area of 1856 m² and is designed to produce up to 450 m³ d⁻¹ at a recovery of 90%. The membranes operate at a mean transmembrane pressure of 2.5 bar producing an average flux of 14 LMH at a temperature of 25–30°C. Membrane cleaning is accomplished by 30 minute cycles of either hot water or caustic soda/proprietary high-pH cleaning agents which are initiated every day and twice a week respectively.

Carbon filter
The final stage of the treatment train is an activated carbon filter with a working capacity of 12.5 m³. The bed is operated at a design empty bed contact time of 30 minutes.

5.5.3 Performance
The plant has been successfully operated since 1995 and has proved the ability of the plant to recover all solid and liquid outputs thus generating near-zero discharge. A mass balance across the process demonstrates the concept (Fig. 5.12) with water losses occurring only in the final product and from the driers in the evaporation stage of production. The RO concentrate contains cellulose and lignins which are returned for use in the resin binder. The filter press produces 480 tonnes y⁻¹ of dry solids which are burnt in the boiler or reused as feedstock at the front end of the chipboard production line as a 45–50% dry solids cake. Some 659 tonnes of water enter the production process with 78% lost during the
drying process and 18% recycled from the effluent treatment plant which goes to make up 60% of the boiler feedwater.

The two main water quality parameters measured through the plant are COD and suspended solids. Some 65% of the COD is removed prior to the membrane plant with the permeate containing 1% of the raw effluent value and the final product water < 1% after carbon polishing. Overall the COD of the plant has been reduced from an influent concentration of 20 000 mg l⁻¹ down to < 200 mg l⁻¹ post activated carbon. Almost all the suspended solids are removed in the filter press with a residual of less than 1 mg l⁻¹ entering the RO stage. Suspended solids in the RO permeates are below limits of detection and the total dissolved solids ~1. The product water quality is very soft with a total hardness concentration of 1 mg l⁻¹ and 0.5 mg l⁻¹ (Ca) (Table 5.20) and contains negligible concentrations of silica or sulphate making the water suitable for reuse in the (low-pressure) boiler house for steam.

The treatment plant was built under a lease-purchase agreement where Kronospan made an initial payment of £200 000 ($310 200) in February 1996 and a final payment of £200 000 ($325 000) in February 1999. Under the contract, Esmil operated the plant for a monthly fee of £22 000 ($33 528; 1998) ensuring the treatment of all wash water effluent to meet Kronospan’s water quality objectives, up to an agreed daily maximum. The plant generates an annual saving of £251 740 ($383 365) of which the majority was generated from the avoidance of tankering the effluent away (91%) (Table 5.7). Actual recycling on the plant produces the remaining savings of which 5.6% come from reduced mains water and 3.4% by recovery of raw material. The payback period for the initial payment was less than 10 months and Esmil continued to operate the plant until autumn of 2002 when Kronospan took over responsibility.

The scheme was the first plant worldwide to apply such an approach to MDF effluent and subsequent plants have been installed across Europe. The scheme at
Table 5.7  Cost sheet for MDF wastewater recycling plant (as of 1998)

<table>
<thead>
<tr>
<th></th>
<th>Annual cost, £ ($)</th>
<th>Annual saving, £ ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supplier operator charge</td>
<td>264 000 (430 320)</td>
<td></td>
</tr>
<tr>
<td>Avoidance of tankers charges</td>
<td></td>
<td>468 500 (763 655)</td>
</tr>
<tr>
<td>Reduced mains water charge</td>
<td>29 480 (48 052)</td>
<td></td>
</tr>
<tr>
<td>Raw material recovery</td>
<td>17 760 (28 948)</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>264 000 (430 320)</td>
<td>515 740 (840 656)</td>
</tr>
</tbody>
</table>


Chirk led in part to Esmil being awarded both the Queen's award for environmental achievement (UK) and the department of trade and industry (DTI, UK) award for best environmental practice.

5.6  M-Real Kirkniemi paper mill (Finland)

5.6.1  Background

M-Real's (formerly Metsa Serla) paper mill at Kirkniemi, Finland, is one of the largest integrated paper mills in Scandinavia producing over 350 000 tons of fine paper, 350 000 tons of magazine paper and its own SGW, PRMP pulp every year. M-Real prides itself on being a leader in forest industry environmental protection and was the first to introduce a biological treatment plant to process wastewater in Finland. In 1994 the white water loop was closed with the integration of membrane technology. As a consequence, freshwater intake at the Kirkniemi mill now stands at a record low level of less than 1400 gallons ton⁻¹ product (5.3 m³ ton⁻¹).

5.6.2  Description of plant

At full capacity the plant processes 1400 gallons min⁻¹ (200 m³ h⁻¹) of which 60–70% is reused for shower waters and dilution water for paper machine chemicals. The remaining 30–40% is further treated and used for warm water replacement. The flow initially passes through a screen to the UF feed tank from where it is pumped to the main UF plant (Fig. 5.13).

The UF plant contains 9 cross-rotational (CR) ultrafiltration membranes arranged in a 5:3:1 array (Fig. 5.14). The membranes are regenerated cellulose, polysulphone stack disc modules supplied by Nadir Filtration GmbH and are rated at a molecule weight cut-off of 30 kDa. The plant contains a total membrane area of 720 m² and is designed to treat 216 m³ h⁻¹ at an equivalent approximate recovery of 90–95%. The membranes operate at an average pressure of 0.8 bar producing an average flux of 300 LMH (range 250–500 LMH).
Figure 5.13 Process flow diagram of Kirkniemi white water recycling plant

at a cross-flow velocity of 13 m s\(^{-1}\). Fouling control in the cross-rotational membrane unit is achieved by rotors between all membrane stacks thus generating shear directly at the membrane’s surface. Membrane cleaning is conducted by a combination of acid and alkaline detergents every 5 days for a duration of 60 minutes. Concentrate from the stacks is combined and sent to the biological treatment plant prior to discharge.

Permeate from the UF plant then is split into two streams with 30–40\% of the flow being sent for further treatment in a nanofiltration plant. The plant contains a total membrane area of 900 m\(^2\) and is designed to process 30–40 m\(^3\) h\(^{-1}\) at a temperature of 58–60°C and an overall recovery of 80\%. The membranes are polyaramide/polysulphone spiral-wound modules and are rated at a salt rejection of 50\%. The membranes operate at a TMP of 9 bar which delivers a mean flux of 30–40 LMH. Membrane cleaning in conducted every 3 days with an alkaline detergent and every week with an acid detergent. The membrane life of the plant is in excess of 2 years.

5.6.3 Performance

The UF plant effectively removes all suspended solids and colloidal materials from the feed (Table 5.20). COD removal is only 15\% but the final water quality is suitable for dilution water for paper machine chemicals. The use of the ultrafiltration plant has not only reduced freshwater intake but has improved maintenance and repair of the paper machines which may be due to the fact that the water is free from suspended solids and microorganisms. The cross-rotational UF plant incurs an energy demand of 1.5 kW m\(^{-3}\) which equates to an operating cost of €0.12 m\(^{-3}\) ($0.12 m\(^{-3}\)).

The white water contains sugars, fatty acids and other wood extracts which all exhibit a high fouling propensity. A key aspect of the scheme has been the identification and development of a membrane system that can effectively operate with such a feed flow. The use of highly hydrophilic, almost uncharged regenerated cellulose membranes has had a significant influence on the efficacy of the technology to this application. Operation of the system has been below the critical flux such that only gradual fouling of dissolved species rather than rapid
fouling of suspended solids occurs. This has resulted in cleaning strategies based on routine maintenance procedures rather than on demand and as such offers the potential for further reductions in opex as confidence grows about the limits of the system.

5.7 T. Forsell and Son: dyewaste water recycling plant (UK)

5.7.1 Background

Thomas Forsell and Son Ltd is a family owned and run textile mill founded in the centre of Leicester in 1850, moving to South Wigston (south of the city) in 1963. The main basis of the company is the production of dyed and spun woollen and worsted yarns for the Ministry of Defence. Due to the need for the final product to retain its colour over prolonged periods the dyes are often highly stable chemical compounds which are poorly biodegradable. The dyes tend have poor fixation properties with as much as 50% of the dye being passed to the effluent. Traditionally, the textile house has discharged the effluent, including the dye, to the sewer for downstream treatment at the local sewage works at South Wigston. Dye removal through typical sewage works is low and so the final effluent typically remains high in colour. In January 1996 the Environment Agency imposed colour consents on Severn Trent discharges to the river Sense at sewage works such as South Wigston STW (Table 5.8). The cost of upgrading South Wigston to comply with the new consents was estimated at £1–2.5 m$^{-3}$ ($1.55–3.87 m$^{-3}$). Consequently, Severn Trent passed these consents on to the
Table 5.8  Colour consents

<table>
<thead>
<tr>
<th>Wavelength (nm)</th>
<th>Colour absorbency</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Consent</td>
</tr>
<tr>
<td>500</td>
<td>&lt;0.057</td>
</tr>
<tr>
<td>550</td>
<td>&lt;0.053</td>
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<tr>
<td>600</td>
<td>&lt;0.047</td>
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<tr>
<td>650</td>
<td>&lt;0.047</td>
</tr>
</tbody>
</table>

individual textile companies including T. Forsell and Son Ltd together with four other discharges in the area.

The textile plant processes 10 Mg of yarn per day producing a total of 400 m$^3$ of effluent. The effluent typically exceeds the colour consent by between 20 and 80% and contains high concentrations of chromium. The cost to meet the consent is sufficiently high that recovery and reuse of the water becomes a viable economic consideration. The company then embarked on a pilot study to establish the most appropriate technology. The criteria outlined were the ability to produce reusable water (based on colour and chrome) and the robustness to withstand the natural fluctuations in water quality. Four options were tested:

- Flocculation
- Biological treatment
- Advanced oxidation
- Reverse osmosis

Flocculation is probably the most popular form of effluent treatment in the textile industry. The system trialled at T. Forsell and Son utilised a synthetic organic clay with a high anion exchange capacity. The system was relatively simple and suitable for colour removal but was unable to reduce the chromium concentration below 3 ppm. The biological system comprised an aerobic reactor coupled with UV and hydrogen peroxide capable of removing 80% of the COD. The plant was relatively inexpensive at £150 000 ($232 650) but was unable to remove chromium. The third option was chemical oxidation with ferrous sulphate, hydrogen peroxide, sulphuric acid and lime. The process effectively removed chromium but retained a high salt concentration in the product water. In all the options the final product water was unsuitable for reuse and so not economically feasible. Ultimately a membrane-based technology was selected as it enabled reuse of the water and hence recovery of the spent capital.

5.7.2 Description of plant

The process train is relatively simple with minimal pretreatment required ahead of the reverse osmosis unit (Fig. 5.15). The flow initially passes through a 20 μm screen prior to being pumped to the RO plant. Concentrate from the process is treated with appropriate chemicals and the generated sludge separated out from
the liquor before being transferred into a 24 hour storage tank. The spent cleaning chemicals are blended with the treated concentrate prior to discharge into the local watercourse. Sludge from the site is stored and tankered away for disposal.

The main treatment stage is two RO streams. The membranes are spiral-wound polysulphone modules supplied by Osmonics. The plant contains a total surface area of 1152 m² and treats a flow of 480 m³ d⁻¹ at a temperature of 60°C. The membranes operate at a mean TMP of 25 bar which delivers a mean flux of 16.5 LMH. The plant is operated for 20 hours per day, 5 days a week and 48 weeks a year. The remaining 4 hours per day are assigned to membrane cleaning.
which consists of a sodium hydroxide soak at pH 10.5 resulting in a spent chemical flow of 40 m$^3$ d$^{-1}$.

The major challenge with the plant was connected to concentrate disposal which had to remove both colour and chromium from the concentrated brine prior to discharge. A number of techniques were explored including electrolytic treatment, chemical oxidation, chemical reduction and flocculation. Both the electrolytic and chemical reduction techniques were not successfully implemented at this site probably due to other contaminants in the flow. Ultimately, the most successful option was the use of chlorine dioxide as a chemical oxidant. The chemical destroys the chromophore and hence decolourises the water whilst simultaneously releasing the Cr(III). Colfloc flocculent was then used to aid the physical separation of the insoluble chromium from the flow. In total the plant produces 20 m$^3$ d$^{-1}$ of concentrate which requires 10 kg d$^{-1}$ of chlorine dioxide (0.5 g l$^{-1}$) to effectively treat it prior to transport to the storage tank and ultimately disposal.

Once the plant was operational and the discharge volume was significantly reduced Forsell and Sons were able to re-negotiate the consent limits based on a reduce flow of 30 m$^3$ d$^{-1}$ (Table 5.9). The changes had considerable impact in terms of COD, Cr and Zn as it greatly reduced the need to treat the concentrate. Most importantly concentrate levels of heavy metals were below the new discharge consents and so no treatment of these components was necessary. The impact on the process was to reduce the chemical demand of the plant by removing the need for a flocculent.

### 5.7.3 Performance

The mill is no longer operational but the wastewater recovery plant was successfully operated for over two years recovering 95% of the water as RO permeate which was then reused within the plant. It produced an overall removal of 80.5% COD and 59% total dissolved solids with complete removal of suspended solids. Colour removal was almost complete with absorbance below

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Original consent</th>
<th>New consent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flow (m$^3$ d$^{-1}$)</td>
<td>227</td>
<td>30</td>
</tr>
<tr>
<td>Rate (m$^3$ h$^{-1}$)</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>pH</td>
<td>6–10</td>
<td>6–10</td>
</tr>
<tr>
<td>SS (mg l$^{-1}$)</td>
<td>1000</td>
<td>700</td>
</tr>
<tr>
<td>COD (mg l$^{-1}$)</td>
<td>3000</td>
<td>9000</td>
</tr>
<tr>
<td>Cr (mg l$^{-1}$)</td>
<td>10</td>
<td>15</td>
</tr>
<tr>
<td>Zn (mg l$^{-1}$)</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Colour at 500 nm</td>
<td>0.057</td>
<td>0.431</td>
</tr>
<tr>
<td>Colour at 550 nm</td>
<td>0.053</td>
<td>0.401</td>
</tr>
<tr>
<td>Colour at 600 nm</td>
<td>0.047</td>
<td>0.356</td>
</tr>
<tr>
<td>Colour at 650 nm</td>
<td>0.047</td>
<td>0.356</td>
</tr>
</tbody>
</table>
0.001 across all the consented wavelengths. Heavy metal removal was also very good with chromium concentration reducing from 0.75 to less than 0.02 mg l\(^{-1}\).

The installation of the plant was required by changes to discharge consents but the ability to recycle water within the plant provided a number of economic advantages to ensure the scheme became financially viable. Major savings were achieved in the cost of softened mains water (by 95%) and the reduction of effluent charges. A less obvious saving was made by the reduction in heat demand achieved by recycling the previously heated water within the process. The capital cost of the schemes was £275 000 ($448 800) in 1998 and the annual net saving on the reuse plant was £142 635 ($232 780) resulting in a payback time of 1.9 years (Table 5.10). The company ceased production in March 2001 at which point the quarterly effluent bill had reduced from £12 481 ($17 723), prior to the reuse plant, down to £640 ($908). Overall the decision to install the plant resulted in a profit to the company of £182 067 ($258 535) over its operational life.

### 5.8 N food plant wastewater treatment and recycling plant (Japan)

#### 5.8.1 Background

Yokosuka city, situated in the Kanagawa Prefecture has no nearby rivers and as such water is supplied from water treatment plants situated on the Sagami river a distance of 30 km away. N company is a producer of frozen foods (such as Chinese and Japanese dishes) and is situated within Yokosuka city and so

<table>
<thead>
<tr>
<th>Table 5.10 Cost sheet for dye bath reuse scheme (as of 1999)</th>
<th>Annual cost, £ ($)</th>
<th>Annual saving, £ ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Membrane replacement</td>
<td>29 714 (47 806)</td>
<td></td>
</tr>
<tr>
<td>Pre-filter opex</td>
<td>1408 (2265)</td>
<td></td>
</tr>
<tr>
<td>Labour</td>
<td>5760 (9267)</td>
<td></td>
</tr>
<tr>
<td>Power</td>
<td>22 464 (36 142)</td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>2661 (4281)</td>
<td></td>
</tr>
<tr>
<td>Cleaning chemical</td>
<td>960 (1544)</td>
<td></td>
</tr>
<tr>
<td>Maintenance</td>
<td>3679 (5919)</td>
<td></td>
</tr>
<tr>
<td>Chlorine dioxide treatment</td>
<td>37 058 (59 622)</td>
<td></td>
</tr>
<tr>
<td>Total opex £ m ( ^{3} )</td>
<td>1.07 (1.72)</td>
<td></td>
</tr>
<tr>
<td>Effluent disposal</td>
<td>60 015 (96 558)</td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>68 874 (119 811)</td>
<td></td>
</tr>
<tr>
<td>Water softening</td>
<td>9316 (14 988)</td>
<td></td>
</tr>
<tr>
<td>Colour charge</td>
<td>56 828 (91 430)</td>
<td></td>
</tr>
<tr>
<td>Water heating</td>
<td>50 307 (80 938)</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>103 705 (166 850)</td>
<td>246 340 (396 336)</td>
</tr>
</tbody>
</table>

Currency conversion \( \times 1.608 \) based on 12 August 1999 figures (http://www.oanda.com/convert/fxhistory (accessed November 2002)).
is constantly looking for ways to reduce its water consumption whilst maintaining product quality. A number of approaches have been considered to reduce water demand:

- Rainwater recycling
- Well water supply
- Refrigerator and subfloor condensation recycling
- Wastewater recycling

The final preference was for wastewater recycling as it provided the most secure source of water available to the plant. Three applications were identified for the water including pre washing of product materials, cooling units and floor washing and boiler feedwater. The projected demand for each application was 40 tonnes day\(^{-1}\) which was anticipated to require 140 tonnes day\(^{-1}\) in total of untreated wastewater to allow for losses during recovery.

5.8.2 Description of plant

Wastewater is collected and treated in pre existing rotating biological contactors before passing to the primary aeration tank and then onto the submerged membrane bioreactor. The permeate is disinfected and stored in the grey water tank from where it is either pumped for direct use in the cooling tower and the washing system or diverted to a reverse osmosis plant for further treatment prior to use as boiler feed water (Fig. 5.17).

Pretreatment
The flow initially passes through a pre-existing RBC unit followed by a drum filter to remove trash components from the sewage and biomass.

MBR
The primary aeration tank (100 m\(^{3}\)) and MBR (50 m\(^{3}\)) are interconnected to provide a hydraulic residence time of 24 hours which equates to a BOD loading rate of 1 kg BOD m\(^{-1}\) d\(^{-1}\). The biomass is periodically wasted to maintain the mixed liquor concentration within the range 8–12 g l\(^{-1}\) which equates to a

![Figure 5.17](image-url)
sludge retention time of 20 days. This range of mixed liquor concentration is well known to provide optimal performance of the MBR system and is necessary to reduce operational problems that occur at either lower or higher values.

The membrane bioreactor plant (Fig. 5.18) consists of a suspended growth bioreactor into which six double-tier membrane modules are submerged providing a total membrane area of 960 m². The membranes are hydrophilic polyethylene hollow fibres rated at a pore size of 0.1 μm. The system is designed to treat 140 m³ d⁻¹ under a constant flux of 5 LMH. Flow through the membranes is generated by a combination of hydraulic head and permeate suction providing a total transmembrane pressure of between 25 and 40 kPa (0.25–0.4 bar) depending on the level of fouling. Fouling control is achieved by a combination of air scour over the membrane fibres and periods of intermittent operation which relaxes the fouled layer and enables the air scour to remove more material. Chemical cleaning is required every six months by an overnight soak in 0.1% sodium hypochlorite and 4% sodium hydroxide solution. The membranes have as yet not required replacement during the 4 years of operation.

Reverse osmosis

The final stage of the treatment train is a reverse osmosis plant arranged in a 2:2:1 array. The membranes are polyamide spiral-wound modules supplied by Toray. The plant contains a total membrane area of 70 m² and operates at a flux of 24 LMH equating to an overall recovery of 67%. Chemical cleaning occurs once a month by means of a chemical feed washing with a solution of ethanol (15%) at pH 12.

5.8.3 Performance

The treatment plant went online in May 1998 and is still being used today to supply both wash water and boiler feed. The MBR plant effectively removes the majority of the organic and suspended pollution found in the wastewater. The BOD is reduced from 2240 mg l⁻¹ in the feed to less than 1 mg l⁻¹ in the MBR

![Figure 5.18](image-url)  (a) Outside view of water tank and (b) MBR at N-food treatment plant
permeate (Table 5.20). Suspended solids and n-hexane extractable compounds are similarly reduced from 1120 and 475 mg l\(^{-1}\) to less than 1 mg l\(^{-1}\) respectively. The final RO stage produces a permeate with a final conductivity of 125 \(\mu\)S cm\(^{-1}\).

Combined water and sewerage charges in the city of Yokosuka are ¥607 m\(^{-3}\) ($4.15 m\(^{-3}\)). The plant operates for 260 days per year and so the installation and operation of the reuse plant results in a direct saving (Table 5.11) of ¥20,338,000 ($139,315) per annum with an additional ¥1,400,000 ($9590) saved through reduced chemical demands on the boiler system. The total annual costs of running the additional treatment technology is ¥5,406,300 ($37,033) resulting in a net annual saving of ¥14,931,700 ($102,282). The cost of the plant is covered by offsetting ¥3,000,000 ($20,550) per year as depreciation on the plant over its projected lifetime yielding a true annual saving of ¥11,931,700 ($81,732).

5.9 Pasfrost vegetable processing plant (Belgium)

5.9.1 Background

Pasfrost, located in Passchendaele, western Belgium, is a vegetable processing company producing 65,000 tonnes of deep-frozen vegetables per year. The plant processes a wide variety of vegetables depending on the season ranging from squashes and leaf crops in the summer to root vegetables in the winter. Process water is used at the site for a range of activities such as washing/rinsing, blanching, steam peeling, caustic peeling and cooling.

Traditionally, water is supplied from groundwater sources that are bacteriologically safe and require no treatment prior to use. However, economic development in the region has led to increasing pressures on the ground water sources resulting in the need to abstract from wells over 300 m deep. The quality

| Table 5.11 Cost sheet for wastewater treatment reuse plant (as of 1998) |
|-----------------------------|-----------------------------|
| Annual cost (Y)             | Annual saving (Y)           |
| MBR energy (120 m\(^{3}\) d\(^{-1}\)) | 925,500 (6,339) |
| MBR membrane (120 m\(^{3}\) d\(^{-1}\)) | 3,600,000 (24,660) |
| RO energy (40 m\(^{3}\) d\(^{-1}\)) | 280,800 (1,923) |
| RO membrane (40 m\(^{3}\) d\(^{-1}\)) | 600,000 (4,110) |
| Total opex (Y m\(^{-3}\)) | 205,8 (1.4) |

Reduced water charges | 9,469,200 (64,864)
Reduced sewage charges | 9,469,200 (64,864)
Reduction in boiler chemical costs | 1,400,000 (9,590)
Total | 5,406,300 (37,033)

Currency conversion (\(\times 0.00685\)) based on 12 August 1998 figures (http://www.oanda.com/convert/fxhistory (accessed November 2002)).
of the groundwater has also deteriorated (Table 5.12) such that the Flemish government has forced companies in the region to severely reduce abstraction volumes.

To maintain production and enable any future expansions requires both conservation and reuse to be considered. Specific items adopted at the Pasfrost site include:

- Partial reuse of wash water for low-grade application which has decreased the specific water use from 5 down to 3.5 m³ tonne⁻¹ of product.
- Steam peeling instead of caustic peeling to reduce the salt concentration in the waste water.
- Anaerobic pretreatment and extension of aerobic treatment producing a more stable effluent suitable for further treatment and reuse.

By reusing treated wastewater the groundwater demand has been reduced by 50%, equating to a specific water consumption of 2 m³ tonne⁻¹ of product. Specific concerns with the requirements of a reclamation plant were the need to handle feedwaters with highly variable organics levels, minimise operational costs and achieve high membrane life expectancy. A core element to the design of the reclamation facility was the ability to deliver a constant quality feed to the advanced treatment processes.

Overall, introduction of the reuse options has reduced the groundwater requirement to 100 000 m³ y⁻¹ with 200 000 m³ y⁻¹ coming from reuse of partially-treated wastewater and 100 000 m³ y⁻¹ from the advanced water treatment plant (Fig. 5.19). Discharge from the site is 100 000 m³ y⁻¹ of which 50% is evaporation and 50% is discharge from the treatment works.

5.9.2 Description of plant

The flow initially passes through a screen and a heat exchanger to raise the average temperature from 20 to 30°C as a pretreatment for the anaerobic stage. The anaerobic reactor is a 5000 m³ UASB operated at a specific loading rate of 3–6 kg COD m⁻³ d⁻¹ enabling a total of 30 000 kg COD to be treated per day. The produced biogas is used for heating the incoming flow and has an equivalent steam rating of 70 tonnes day⁻¹ based on the production of 5000–6000 m³ d⁻¹ of biogas during high loads.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Groundwater</th>
<th>WHO guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>8.3</td>
<td>6.5–8.5</td>
</tr>
<tr>
<td>SO₄²⁻ (mg L⁻¹)</td>
<td>126</td>
<td>400</td>
</tr>
<tr>
<td>HCO₃⁻ (mg L⁻¹)</td>
<td>552</td>
<td></td>
</tr>
<tr>
<td>Cl⁻ (mg L⁻¹)</td>
<td>550</td>
<td>250</td>
</tr>
<tr>
<td>Conductivity (mS cm⁻¹)</td>
<td>2.6</td>
<td>0.25</td>
</tr>
<tr>
<td>TH (mmol L⁻¹)</td>
<td>0.3</td>
<td></td>
</tr>
</tbody>
</table>
The flow is then passed to a 11,000 m$^3$ activated sludge plant operated at a specific loading rate of 0.1 kg COD kg VSS$^{-1}$ day$^{-1}$ to polish and re-aerate the flow. After sedimentation the effluent is treated in two serial steps by ASTRASAND$^u$ up flow monomedia continuous depth filters. The first set of filters act as a roughing stage and comprise two 5 m$^2$ filter beds operating at a hydraulic filtration velocity of 10 m$^3$ m$^{-2}$ h$^{-1}$. The second stage incorporates a polymer dose to help flocculate and capture the remaining solids as they pass through two 10 m$^2$ filters.

Flow is either directly pumped after chlorination for second-grade operations such as machine washing and cooling or is pumped to the membrane filtration treatment plant. The first membrane stage is a semi dead-end hollow fibre ultrafiltration membrane plant supplied by Norit X-flow designed to deliver an overall flow of 40 m$^3$ h$^{-1}$ (Fig. 5.21). The hydrophilic membranes are manufactured out of PES/PVP blend and have a pore size rating of 50 kDa. The membranes operate over a pressure range of 0.3 to 1 bar which delivers a mean flux of 45 LMH. Cleaning of the membranes occurs by a periodic backflush of permeate and a regular chemical clean.

The final stage of treatment is a two-stage reverse osmosis plant containing spiral-wound neutrally-charged polyamide LFC$^1$ membranes supplied by Hydraunautics. The plant is designed to deliver a net production of 20 m$^3$ h$^{-1}$ at a recovery of 70%. The membranes operate over a pressure range of 8–10 bar, delivering a mean flux of 30 LMH. As an extra precaution the permeate is passed through a UV plant to ensure the water is sterile. The final effluent is then mixed with the ground water and pumped into the production process. RO concentrate is discharged with the excess flow.

5.9.3 Performance

The plant has been operated since mid-2000 and has suffered no overall problems with effluent quality. Incoming wastewater is ultimately reduced in
COD concentration from 12,000 mg l\(^{-1}\) to a non-detectable level by the treatment train (Table 5.13). Final product water quality also shows non-detectable levels of turbidity and coliform concentrations. The recycle water contains a lower concentration of conductivity, iron and total hardness but is slightly higher in NH\(_4^+\)-N and HCO\(_3^-\) compared to the existing groundwater.

The main operational difficulty has been in maintaining flux rates through the UF plant throughout the year. During winter periods (vegetables and Schorseneer) UF fluxes deteriorate to 50% of the design flow. Diagnostics of the membrane surface reveal spots attributable to a variety of humic acids. Chemical cleaning protocols have been changed to use enzymatic cleaners but the overall problem remains. During the winter period the water consumption is reduced and so it was decided to operate at reduced fluxes, which seem to ameliorate the problem. In the summer the UF fluxes can be increased to 50–60 LMH and as a consequence RO capacity has been increased by 40% to 30 m\(^3\) h\(^{-1}\) without additional investment in UF plant.

Operation during the initial phase of the project has yielded a net plant flow of 20 m\(^3\) h\(^{-1}\) while the infrastructure is designed to produce 40 m\(^3\) h\(^{-1}\) and so lost
Table 5.13  Typical water quality profile through the Pasfrost treatment plant

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Wastewater</th>
<th>Post anaerobic</th>
<th>Post aerobic</th>
<th>Post filters</th>
<th>RO/UV</th>
<th>Groundwater</th>
</tr>
</thead>
<tbody>
<tr>
<td>COD (mg l⁻¹)</td>
<td>12 000</td>
<td>1200</td>
<td>114</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>pH</td>
<td>8.5</td>
<td>7</td>
<td>8.3</td>
<td>8.2</td>
<td>5.5</td>
<td>8.3</td>
</tr>
<tr>
<td>Total P (mg l⁻¹)</td>
<td>34</td>
<td>34</td>
<td>&lt;0.5</td>
<td>&lt;0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NH₄⁺-N (mg l⁻¹)</td>
<td>&lt;2</td>
<td>&lt;2</td>
<td>&lt;2</td>
<td>0.17</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe (mg l⁻¹)</td>
<td>0.03</td>
<td>0.23</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TH (mmol l⁻¹)</td>
<td>1</td>
<td>1</td>
<td>&gt;0.3</td>
<td>0.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HCO₃⁻ (g l⁻¹)</td>
<td>2.9</td>
<td>2.9</td>
<td>&lt;0.3</td>
<td>&lt;0.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conductivity (mS cm⁻¹)</td>
<td>4.3</td>
<td>4.3</td>
<td>0.15</td>
<td>2.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Turbidity (NTU)</td>
<td>16</td>
<td>4.3</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CFU (ml⁻¹)</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E-coli (ml⁻¹)</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 5.14  Balance of operating costs at the Pasfrost site

<table>
<thead>
<tr>
<th>Cost</th>
<th>Design, € m⁻¹ ($ m⁻¹)</th>
<th>Actual, € m⁻¹ ($ m⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depreciation on investment</td>
<td>0.49 (0.48)</td>
<td>0.49 (0.48)</td>
</tr>
<tr>
<td>Depreciation on membranes</td>
<td>0.14 (0.14)</td>
<td>0.14 (0.14)</td>
</tr>
<tr>
<td>Chemicals</td>
<td>0.17 (0.17)</td>
<td>0.07 (0.07)</td>
</tr>
<tr>
<td>Energy</td>
<td>0.07 (0.07)</td>
<td>0.03 (0.03)</td>
</tr>
<tr>
<td>Maintenance/operation</td>
<td>0.08 (0.08)</td>
<td>0.06 (0.06)</td>
</tr>
<tr>
<td>Total</td>
<td>0.95 (0.947)</td>
<td>0.79 (0.787)</td>
</tr>
</tbody>
</table>

Currency conversion (× 0.99739) based on 12 August 2002 figures (http://www.oanda.com/convert/fchistory (accessed November 2002)).

capital is expected. The operating costs of the plant during Phase One has been evaluated after two years of operation (Table 5.14) with a total opex of €0.79 m⁻³ ($0.78 m⁻³) including plant depreciation and €0.16 m⁻³ ($0.16 m⁻³) without. The actual opex are lower, if compared to design predictions, which is due to the introduction of steam peeling which has reduced the need for pH correction. In fact during some periods of the year no pH correction is required at all. The other factor has been improved stability of UF operation whilst the TMP remains below 0.5 bar. Annual production is currently 150 000 m³ which represents a saving of €24 000 ($23 900) compared to the original design predictions.

The introduction of the reuse scheme enabled Pasfrost to further extend its production volume, without increasing the specific water costs. As the legislation for groundwater abstraction becomes more stringent the focus will be on increasing the water reuse volume to even higher levels (60–70% of intake). Without the reuse scheme the present production volume would only have been possible if drinking water was used, which needs to be softened on-site. Taking into account the extra drinking water intake and the extra waste water discharge volume, this would result in an additional cost of €0.30–0.40 per m³ in comparison to the actual present specific costs.
5.10 Automotive water recycling (Germany)

5.10.1 Background

The automotive industry requires large volumes of water and chemicals in the production of finished cars and trucks. The majority of the water is associated with the pretreatment and electrocoating stages where the car bodies need to be cleaned prior to the different stages of production, and includes a rinse between each step.

The application of membrane technology within the electrocoating process is widespread. In fact, the process would not be viable without ultrafiltration which has been used for more than 20 years to extract rinsing solutions from paint. The integration of the technology has numerous drivers such as a need to remove drag out paint before stoving, almost total recovery of the paint and the avoidance of effluent problems. In part this has been integral to the development of electropainting techniques and so has become part of the core process rather than additional technology that needs to be justified. This places the industry in a different position from many of the others which are currently considering membrane technology. However, uptake of membranes to other parts of the pretreatment and electrocoating stages has been considerably slower. The result is that still heavy water demands are placed on production with up to $500 \text{ m}^3$ of process water being required per car. In more recent times economic drivers have required the industry to examine the potential to reduce costs of both effluent treatment and chemicals consumption. Two applications where this is becoming more established is in removal of oil from the pretreatment cleaners at the start of the production and recycling and recovery of paint from the final rinse water at the end. The economic benefits are to a large extent country-specific due to differences in the available water and the disposal options for the effluent. However, the post-paint rinse water offers the most obvious benefits due to the recovery of high-value paint and so will be examined further.

Once the electrophoretic painting is complete drag out is removed from the car bodies by rinsing with ultrafiltration permeate extracted from the paint. A final deionised water rinse is then applied to remove final paint traces and salts which may otherwise reduce product quality (Fig. 5.22). The wastewater produced from this process contains paint solids which have to be treated in the liquid effluent line prior to discharge and the solids disposed as a contaminated waste. A number of techniques have been tried to reduce demand on water and effluent and to recover lost paint:

1. Extension of UF rinsing
2. Extraction of RO permeate from UF
3. Treatment of post-paint rinse water by UF

**Extension** of using UF permeate for the entire rinse period and thus replacing the deionised water (option 1) potentially has shortcomings as the permeate still
retains salts and other soluble impurities. Potential stability problems could arise when subsequent paint layers are coated, resulting in a poor quality finish. The impurities can be removed by utilising reverse osmosis membranes to further purify the rinse water (option 2). However, the feed to the RO system is likely to have a high fouling propensity making the option technically undesirable. In both options, additional UF permeate is required and this is a relatively expensive option since less paint is recovered than in the primary UF rinsing. Ultimately, a separate recycling loop was selected which involves recycling of the post-paint rinse water and recovery of paint (option 3). An additional benefit of option 3 is that a small concentration of solvent builds up within the loop which improves the efficacy of the rinsing solution over that of deionised water.

5.10.2 Plant description

The treatment train comprises a treatment tank, a recirculation pump and an ultrafiltration module (Fig. 5.23). Used water-containing paint particles are retained by the membrane and concentrated in the treatment tank. When the concentration in the treatment tank is sufficient the contents are pumped to the electrocoat tank replacing make-up water. This is important to the success of the scheme as it represents a recovery of valuable paint product.

The ultrafiltration modules are acrylonitrile plate-and-frame membranes (Fig. 2.7) supplied by Rhodia Orelis rated at a molecular weight cut off of 50 kDa. The plant contains a total membrane area of 55 m² and is designed to treat a flow of 192 m³ d⁻¹ at a temperature of 40°C. The membranes are operated over a TMP range of 1–3 bar delivering a flux range of 145–300 LMH at a cross flow velocity of 2.6 m s⁻¹. Cleaning occurs approximately every 3–4 months and involves a 300-minute cleaning cycle with organic acid and solvents. The operation of the
plant results in a membrane replacement life of over 3 years. Membrane integrity is checked during both manufacture and operation. During manufacture membrane quality is monitored using air permeability tests. During operation, permeate quality is monitored by a combination of visual inspection and on-line turbidity meters. Any individual permeate tubes seen to be passing retentate are removed from the collector and diverted back to the treatment tank and so there is no need to make interventions for incidental damage.

5.10.3 **Performance**

The plant operates within the required specifications for recycling water and recovering paint. The main operational concern with the system is the stability of the paint solids which would otherwise coat the membrane, a problem which is controlled by pH adjustment. Bacteria contamination is also a problem due to the close contact of the rinse water with air. Growth is controlled through the use of paint-compatible biocide, but ultimately the membranes may need mechanically cleaning by hand.

The high water demand of the process and the ability to recover valuable paint make the recycling scheme very favourable economically. The plant reduces the requirement for water and effluent treatment by 60 000 m\(^3\) y\(^{-1}\) resulting in a €327 000 ($326 146) saving. Some 18 600 kg y\(^{-1}\) of paint are also recovered representing 22% of the total annual saving accrued from the plant. The remaining saving are in labour costs associated with having to clean the rinse water tank less regularly. The major cost of running the plant is energy to run the pumps and cooling circuit which equates to a total annual energy demand of 324 000 kWh. The remaining costs are due to membrane cleaning and replacement resulting in a total opex of €0.36 m\(^{-3}\) ($0.355 m\(^{-3}\)). The plant cost a capital outlay of €270 000 ($269 000) with a further €24 000 ($23 600) required for first year financing. The annual net cash flow is €435 000 ($433 900) resulting in a pay back period of less than 8 months. Although the economic success of the plant is country-specific a similar scheme in the UK
Table 5.15  Cost sheet for post paint rinse water reuse plant (as of 2002)

<table>
<thead>
<tr>
<th></th>
<th>Annual cost, € ($)</th>
<th>Annual saving, € ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy</td>
<td>16 500 (16 456)</td>
<td></td>
</tr>
<tr>
<td>Membrane cleaning</td>
<td>5500 (5485)</td>
<td></td>
</tr>
<tr>
<td>Recovered paint</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water and effluent</td>
<td>327 000 (326 146)</td>
<td></td>
</tr>
<tr>
<td>Labour</td>
<td>30 000 (29 921)</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>22 000 (21 942)</td>
<td>457 000 (455 807)</td>
</tr>
</tbody>
</table>

Currency conversion (× 0.99739) based on 12 August 2002 figures (http://www.oanda.com/convert/fxhistory (accessed November 2002)).

would have a payback period in the region of 13 months due to the reduced water prices.

5.11 NEC Semiconductors: microelectronics wastewater reclamation (UK)

5.11.1 Background

NEC Semiconductors (UK) Ltd, Livingston, was formed in 1982 as part of the NEC Corporation and was originally assigned for assembly and testing before expanding into manufacture. The fabrication facility produces several millions chips per month and has the largest operational clean room floor area in Europe (dated 1999). NEC incorporates reclamation and reuse within its business strategy achieving ISO 14001 accreditation in December 1991. Included in the strategy is both reduced chemical consumption and waste recycling. For instance, reduction in machine bath volume and photo-resist dispensing volumes generated significant savings in hydrofluoric acid (£40 000 ($62 040)). Materials such as cardboard, paper, plastics, acids and reject silicon wafers are also recycled. For instance, the waste silicon is used in aluminium production acting as a strengthening agent for the final product.

The initial driver for water recycling was to ensure sufficient water resources at facilities where external water was limited. The experience gained at these sites has led to reclaim plants operating at sites where water is plentiful and inexpensive. In such cases the driver for reuse is to reduce operating and capital costs of water supply with the added benefits of environmental preservation and associated publicity. Reclamation at some sites has even extended to ZLD (zero liquid discharge).

The majority of the DI water required at the production facility is used in the wet bench machines in the clean room where the silicon wafers pass through at various stages during production. The wet benches consist of a concentrated acid bath, for etching, followed by a series of rinse baths. The wafers are initially lowered into the acid bath for a preset time to achieve the required etch depth.
Afterwards the wafers are placed in a constantly overflowing deionised (DI) water bath and then onto a second rinse stage before moving onto the next stage of production. The DI water can be either hot or cold, depending on the acid bath temperature, and is drained separately from the bench before being pumped to the treatment facility.

Transistor gate sizes can be smaller than 2 μm and as such any ions or the particles remaining on the silicon wafers can cause short circuits. Consequently the DI water quality standards need to ensure high-purity water (Table 5.16).

Raw water supply to the production facility is high in organics due to the moorland intake and highly variable due to limited treatment at the local water treatment works. Consequently the potable water intake requires a large number of treatment processes prior to entering the DI production facility (Fig. 5.24). In fact, the front end of the works is similar to an advanced potable water works with coagulation-DAF and dual media depth filters to remove solids and activated carbon to remove organics and chlorine. Following the GAC is an anionic organic scavenger resin bed, a cartridge filter and then finally a reverse osmosis plant. The efficacy of the GAC in removing chlorine is essential to protect the resin and RO membrane from oxidation. The organic scavenger resin is regenerated on site with brine and caustic solution and contains two streams allowing for maintenance and regeneration.

The water then enters the DI plant which is split into primary and polishing stages. In the primary stage the water passes through a sequence of cation–anion–cation ion exchange beds, a 10 μm cartridge filter to remove resin and precipitated organics and a 254 nm UV plant. Following the UV stage the water is filtered through a 3-stage RO plant arranged in a 7:3:2 array. The permeate is de-aerated prior to being pumped to the polishing stage of production. In the final stage the water passes through a cooler, 185 nm UV and UF membrane filtration plant before being put into the supply ring main. Total production of ultra pure water is 200 m³ h⁻¹ at an overall recovery rate of 71%.

In comparison to the intake raw water the DI waste water is very low in organics and solids and so requires relatively less treatment to produce a water of sufficient quality to enter the primary DI production stage. The main difference between the water sources is the very low cation and bacteria count in the reclaim water (Table 5.17). This is to be expected as the reclaim water is generated from the DI baths, which follow high-purity acid baths. Importantly,

Table 5.16 Water quality standards for DI water

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resistivity</td>
<td>18 MΩ cm⁻²</td>
</tr>
<tr>
<td>Particles (&gt; 0.05 μm)</td>
<td>&lt; 5 ml⁻¹</td>
</tr>
<tr>
<td>Bacteria</td>
<td>&lt; 1 μl⁻¹</td>
</tr>
<tr>
<td>TOC</td>
<td>&lt; 2 μg l⁻¹</td>
</tr>
<tr>
<td>DO</td>
<td>&lt; 50 μg l⁻¹</td>
</tr>
<tr>
<td>Metals</td>
<td>&lt; 0.01 μg l⁻¹</td>
</tr>
</tbody>
</table>
as the baths are situated in the clean rooms no contamination is picked up from the atmosphere. The reclaim water does, however, contain much higher concentrations of sulphate from sulphuric acid remaining on the wafers.

An important aspect of the scheme is that the use of reclaimed water does not affect product quality in any way. To ensure this, reclaimed water is monitored in terms of TOC and conductivity prior to being pumped to the primary DI production stage. Any waste above the allowable levels is rejected and passes to the effluent treatment plant. The plant was originally designed to accept water below 300 µS cm⁻¹ but has had to increase this to 600 µS cm⁻¹ to increase the amount of water through the reclaim plant.

5.11.2 Description of plant

The reclaim water is essentially deionised water with sulphate anions added; no cation removal is therefore required and the process flow sheet reflects this (Fig. 5.25). Reclaim water is pumped at a rate of 140 m³ d⁻¹ through a 14 m³ activated carbon bed (diameter = 2.8 m, height = 3.05 m) for the removal of hydrogen peroxide which can be present in some of the baths. This is necessary to protect the downstream processes from oxidation. After the adsorption bed the water passes through a 2.2 m diameter weak anion exchange bed with a WBV of 5.2 m³. The anion bed removes free ions of sulphate, phosphate and fluoride. The anion resin is regenerated with a caustic solution after a preset number of bed volumes have been treated or the pH falls below pH 6.

The water then passes through a 10 µm filter to remove any resin beads or activated carbon particles. An UV lamp operating at a wavelength of 185 nm
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Reclaim water</th>
<th>Raw water</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min.</td>
<td>Max.</td>
</tr>
<tr>
<td>Bacteria (cells l⁻¹)</td>
<td>5</td>
<td>735</td>
</tr>
<tr>
<td>Conductivity (µS cm⁻¹)</td>
<td>250</td>
<td>600</td>
</tr>
<tr>
<td>Silica (mg l⁻¹)</td>
<td>0.5</td>
<td>32</td>
</tr>
<tr>
<td>Iron (mg l⁻¹)</td>
<td>0.01</td>
<td>0.1</td>
</tr>
<tr>
<td>Alkalinity (mg l⁻¹)</td>
<td>0.004</td>
<td>4.9</td>
</tr>
<tr>
<td>Calcium (mg l⁻¹)</td>
<td>0.01</td>
<td>0.67</td>
</tr>
<tr>
<td>Magnesium (mg l⁻¹)</td>
<td>0.01</td>
<td>0.07</td>
</tr>
<tr>
<td>Chloride (mg l⁻¹)</td>
<td>0.02</td>
<td>19.8</td>
</tr>
<tr>
<td>Phosphate (mg l⁻¹)</td>
<td>0.05</td>
<td>2.05</td>
</tr>
<tr>
<td>Sulphate (mg l⁻¹)</td>
<td>30</td>
<td>110</td>
</tr>
<tr>
<td>Nitrate (mg l⁻¹)</td>
<td>0.13</td>
<td>1.2</td>
</tr>
<tr>
<td>Fluoride (mg l⁻¹)</td>
<td>1</td>
<td>15.5</td>
</tr>
</tbody>
</table>

![Figure 5.25](image_url)  
*Figure 5.25 Process flow diagram of the reclaim treatment facility at NEC Semiconductors Ltd*

then sterilises the water and removes any trace organics remaining in the flow. The final stage of the treatment is a reverse osmosis plant consisting of 8 inch × 8 inch modules each fitted with four elements and arranged in a 5:2:1 array. The plant operates at a 90% recovery generating 130 m³ h⁻¹ of treated reclaim water.

### 5.11.3 Performance

Water enters the reclaim plant with a conductivity of 600 µS cm⁻¹. The weak anion bed reduces the conductivity to 25 µS cm⁻¹ which is then further reduced through the RO plant to a final value of 3 µS cm⁻¹. The water then mixes with the treated raw water before being passed onto the DI production facility where
water is ultimately produced with a resistivity of 18.2 $\text{M}\Omega\text{ cm}^{-2}$. Some 58% of the high-purity water is reclaimed and recycled to the ultrapure water plant. The remaining water is sent to the effluent treatment facility with the RO concentrate and the brine regenerate. Under these conditions 54% of the production flow is generated from recycled flow.

Included in the reclamation strategy is the utilisation of waste heat through a heat exchange network. Hot reclaim drains are used to preheat the supply of high purity DI. This represents a huge energy saving by increasing the cold DI from 25 to 55°C, and has a dual benefit as it cools the wastewater prior to being treated in the reclaim plant.

There is a significant financial saving from using reclaim water as 83% of the treatment cost is associated with bringing raw water on site and then disposing of it (Table 5.18). The reclaim water costs only 12% that of treating raw water generating a saving of 20% on the total operating and maintenance budget for the DI water treatment plant. The main economic decision is concerned with determining the amount of reclaim water to recycle. Zero reclaim incurs high operational costs due to raw water price but 100% reclaim incurs excessive additional up front capital investment. Between the two extremes is an economic optimum, which depends largely on the utility cost of the local region (Fig. 5.26). In order for the reclaim percentage to increase the cost curve needs to change such that the capital cost in relation to the raw water/discharge costs will decrease. The major barrier to this is the rapid changes that can occur in microprocessor production which means new chemical pollutant can be generated which the original plant is incapable of treating.

### 5.12 Discussion

The case studies featured have not only demonstrated that reuse of industrial waste water is technically feasible but economically viable. In all cases the installation of a membrane system for recycling water has resulted in an overall saving to the company involved. Annual savings have ranged from $102,282 to $642,240 (Table 5.19) and have paid back the initial capital from within 8 months (automotive) to 6 years (power). The most cost-effective of the schemes has been the automotive plant (internal loop) which generates an annual return

#### Table 5.18 Water treatment cost breakdown

<table>
<thead>
<tr>
<th></th>
<th>Raw water (%)</th>
<th>Reclaim water (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raw water (%)</td>
<td>67.2</td>
<td>0</td>
</tr>
<tr>
<td>Electricity (%)</td>
<td>13.4</td>
<td>9.41</td>
</tr>
<tr>
<td>Chemicals (%)</td>
<td>3.2</td>
<td>2.28</td>
</tr>
<tr>
<td>Discharge (%)</td>
<td>16.1</td>
<td>0</td>
</tr>
<tr>
<td>Total (%)</td>
<td>100</td>
<td>11.7</td>
</tr>
</tbody>
</table>

Excludes labour and media replacement.
on investment of 147% equating to a profit of $2259 m\textsuperscript{3} d\textsuperscript{-1}. The least profitable of the schemes has been the power station at Eraring (external loop) which generates an annual return on investment of 19.2% equating to a profit rate of $171 m\textsuperscript{3} d\textsuperscript{-1}.

The savings have been generated from a number of sources in addition to the direct benefit of using less external (potable) water. In the case of the power stations, recycling has decreased the number of regenerations required in the demineraliser plant either reducing operating costs (Eraring) and/or increasing capacity (Flag Fen). In other cases recovery of raw materials has increased the profit margins, such as paint (Germany) and pulp solids (Chirk). Perhaps a less obvious saving has been in energy savings by recycling hot water streams and so reducing the heat requirement at the site (Livingston, South Wigston, Apeldoorn).

However, the driver for reuse is not always directly a financial one. In the majority of case studies outlined here the original interest in reclamation was

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**Table 5.19 Summary of case studies**

<table>
<thead>
<tr>
<th>Industry</th>
<th>Flow rate (m\textsuperscript{3} d\textsuperscript{-1})</th>
<th>Specific flux (LMH bar\textsuperscript{1})</th>
<th>Capital investment ($)</th>
<th>Annual saving ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power (2000)</td>
<td>1200</td>
<td>3.63 (RO)</td>
<td>1524000</td>
<td>–</td>
</tr>
<tr>
<td>Power (1994)</td>
<td>3750</td>
<td>0.88 (RO)</td>
<td>3340000</td>
<td>642240</td>
</tr>
<tr>
<td>Power (1991)</td>
<td>1363</td>
<td>–</td>
<td>750000</td>
<td>240000</td>
</tr>
<tr>
<td>Paper (2001)</td>
<td>216</td>
<td>34.3 (UF)</td>
<td>–</td>
<td>226391\textsuperscript{a}</td>
</tr>
<tr>
<td>MDF (1995)</td>
<td>400</td>
<td>0.56 (RO)</td>
<td>635200</td>
<td>383365</td>
</tr>
<tr>
<td>Paper (1994)</td>
<td>5184</td>
<td>375 (UF)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Textile (1996)</td>
<td>480</td>
<td>0.66 (RO)</td>
<td>448635</td>
<td>232780</td>
</tr>
<tr>
<td>Food (1996)</td>
<td>140</td>
<td>15.4 (MF)</td>
<td>–</td>
<td>102282</td>
</tr>
<tr>
<td>Food (2000)</td>
<td>685</td>
<td>3.3 (RO)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Automotive (2000)</td>
<td>192</td>
<td>111 (UF)</td>
<td>293223</td>
<td>433864</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Data in scheme not complete.
generated from an indirect financial driver such as changes to legislation (textile, food) or a need to secure sufficient water supplies. For instance, in the case of the textile plant, legislation imposed a treatment requirement on the plant. Once money had to be spent to comply with the legislation, the benefits of ensuring the water could be recycled became important as otherwise the capital expenditure could not be recovered. The regulation requirements can go as far as zero liquid discharge, in which case reuse is a necessity rather than an option. In such cases, the driver is to reduce overall treatment costs (Doswell). However, in a number of the schemes the driver has been directly one of reducing operating costs of which water supply can be a major component (Eraring, Livingston).

The selection of membranes in the reclamation process train has occurred in a number of ways. In some industries either membrane technology is already used (Germany, Livingston) or the plant involves similar levels of technology (Eraring). At other schemes the use of membranes has been a radically new development (South Wigston). The familiarity with the technology appears to be in part linked to the need for indirect financial drivers to exist before reuse is considered. This is probably because a high degree of confidence is required in the technology designed for reclaiming the water, since it cannot be allowed to adversely affect core production quality.

As expected it is difficult to draw commonalities from a broad range of industries. However, a number of points can be concluded. The key facet of the reclamation system in all cases has been its ability to withstand variations in the wastewater quality, whilst producing a water quality suitable for reuse. Effluent quality from membranes is usually very good, such that the main concern is achieving sufficient throughput without incurring excessive cost. This is reflected in the level of pretreatment required from the different schemes. In cases where simple closed loops are being generated the pretreatment requirement is minimal (as in the automotive industry). However, in situations such as the reuse of secondary effluent and other wastes with high fouling propensities, more involved pretreatment is required. In some cases this involves two membrane stages and in other cases more traditional pretreatment (such as coagulation followed by depth filtration).

Overall, the case studies have shown the suitability of membrane technologies in particular for industrial effluent recovery and reuse. The ability to produce reclaimed water of sufficient quality is clear. However, the throughputs are quite different between the schemes. For instance, comparing the specific fluxes of the four RO schemes described reveals a range between 0.56 and 3.63 LMH bar⁻¹ reflecting the differences in the RO feed water matrices. This demonstrates that each scheme is in part unique, potentially involving problems that have not been encountered in other industries. Moreover a common problem with potential industrial reuse schemes is a paucity of data describing the water quality and hence the design limits. Table 5.20 clearly illustrates this point where in some cases little or no water quality data are known in actual operating schemes. Although easily remedied, data paucity remains a major barrier to uptake of not only membranes for reuse but any treatment technology.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Application</th>
<th>Power</th>
<th>Power</th>
<th>Power</th>
<th>Pulp+ paper</th>
<th>MDF</th>
<th>Pulp+ paper</th>
<th>Textile</th>
<th>Food</th>
<th>Food</th>
<th>Automotive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name of installation</td>
<td>Flag Fen (MF)</td>
<td>Flag Fen (RO)</td>
<td>Eraring</td>
<td>Alpheus</td>
<td>Apeldoorn</td>
<td>Chirk</td>
<td>Kirkniemi</td>
<td>South</td>
<td>Wigston</td>
<td>Wigan</td>
<td>Vitality</td>
</tr>
<tr>
<td>Company</td>
<td>Alpheus</td>
<td>Vivendi</td>
<td>Triqua BV</td>
<td>Esmil Process Systems</td>
<td>Esmil</td>
<td>Systems</td>
<td>Esmil Membranes</td>
<td>Kanagawa</td>
<td>Passchendaele</td>
<td>Germany</td>
<td></td>
</tr>
<tr>
<td>Membrane geometry</td>
<td>Hollow fibre</td>
<td>Spiral wound</td>
<td>Spiral wound</td>
<td>Tube</td>
<td>Spiral wound</td>
<td>CR filter (flat sheet)</td>
<td>Spiral wound</td>
<td>Tomato</td>
<td>Paques</td>
<td>Rhodia</td>
<td></td>
</tr>
<tr>
<td>Height (mm)</td>
<td>2000</td>
<td>1524</td>
<td>1016</td>
<td>1000</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>486</td>
<td>2600</td>
<td>900</td>
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<td>Width (mm)</td>
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<td>48</td>
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<td>310</td>
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<td>Diameter (mm)</td>
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<td>200</td>
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<td>798</td>
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<tr>
<td>Supplier</td>
<td>PALL filtration</td>
<td>Koch</td>
<td>-</td>
<td>-</td>
<td>Osmonics</td>
<td>Nadir Filtration GmbH</td>
<td>PCI Membranes</td>
<td>960</td>
<td>700</td>
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<tr>
<td>Membrane area (m²)</td>
<td>1700</td>
<td>1451</td>
<td>-</td>
<td>-</td>
<td>31.6</td>
<td>80</td>
<td>31.6</td>
<td>Polyethylene</td>
<td>160</td>
<td>35</td>
<td>55</td>
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<tr>
<td>Surface area per module (m²)</td>
<td>50</td>
<td>52</td>
<td>-</td>
<td>-</td>
<td>31.6</td>
<td>80</td>
<td>31.6</td>
<td>Polyethylene</td>
<td>160</td>
<td>35</td>
<td>55</td>
</tr>
<tr>
<td>Material</td>
<td>PVDF</td>
<td>Polyamide</td>
<td>Cellulose acetate</td>
<td>PVDF</td>
<td>Polyamid</td>
<td>Regenerated cellulose, polysulphone</td>
<td>Polysulphone</td>
<td>Polyethylene</td>
<td>PFS/PVP</td>
<td>Acrylonitrile</td>
<td></td>
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<tr>
<td>Pore size (μm)</td>
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<td>-</td>
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<td>0.04</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Material cut-off (kDa)</td>
<td>-</td>
<td>200</td>
<td>98% salt rejection</td>
<td>-</td>
<td>30-100</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>50</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>Surface charge</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Anionic</td>
<td>Slightly negative</td>
<td>-</td>
<td>-</td>
<td>Anionic</td>
<td>Hydrophilic</td>
<td>Hydrophilic</td>
<td>Positive</td>
</tr>
<tr>
<td>Hydrophobicity</td>
<td>Hydrophilic</td>
<td>Hydrophobic</td>
<td>Hydrophilic</td>
<td>Hydrophilic</td>
<td>Hydrophilic</td>
<td>Hydrophilic</td>
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<td>Power</td>
<td>Power</td>
<td>Pulp+</td>
<td>MDF</td>
<td>Pulp+</td>
<td>Textile</td>
<td>Food</td>
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<td>Flag Fen (RO)</td>
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<td>Apeldoorn</td>
<td>Chirk</td>
<td>Kirkniemi</td>
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<td>6 months</td>
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<td>6 months</td>
<td>Variable</td>
<td>3 4 months</td>
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<td>Cleaning chemicals</td>
<td>Hypochlorite+</td>
<td>High + low</td>
<td>Anionic/ionic</td>
<td>Caustic soda, proprietary</td>
<td>Alkaline and acid detergent for</td>
<td>Sodium hypochlorite+</td>
<td>HCl + NaOCl</td>
<td>Organic acid+</td>
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<td>pH</td>
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<td>Flow per module (m³ h⁻¹)</td>
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<td>13</td>
<td>15</td>
<td>1.5</td>
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<td>2</td>
<td>&gt;5</td>
<td>&gt;3</td>
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<td>Cross-flow velocity (m s⁻¹)</td>
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<td>Membrane replacement (years)</td>
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(Table continued on next page)
Table 20B (continued)

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<th>Parameter</th>
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<th>Power</th>
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<th>MDF</th>
<th>Pulp+ paper</th>
<th>Textile</th>
<th>Food</th>
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<td>Power consumption (kW m⁻¹)</td>
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<td>Operating costs ($ m⁻¹)</td>
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<td>Competitive</td>
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LMH = l m⁻³ h⁻¹.
### Table 5.20C Performance

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<th>Power</th>
<th>Power</th>
<th>Pulp + paper</th>
<th>MDF</th>
<th>Pulp + paper</th>
<th>Textile</th>
<th>Food</th>
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<td>COD$_{\text{in-out}}$ (ppm)</td>
<td>Flag Fen (MF)</td>
<td>51.3</td>
<td>-</td>
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<td>-</td>
<td>8000 - 200</td>
<td>15%</td>
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<td>BOD$_{\text{in-out}}$ (ppm)</td>
<td>Flag Fen (RO)</td>
<td>10</td>
<td>20 - 5</td>
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<td>2240 - 1</td>
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<td>Total N$_{\text{in-out}}$ (ppm)</td>
<td>Eraring</td>
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<td>Total P$_{\text{in-out}}$ (ppm)</td>
<td>Apeldoorn</td>
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<td>SS$_{\text{in-out}}$ (ppm)</td>
<td>South Wigston</td>
<td>16 - 2</td>
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<td>Turbidity$_{\text{in-out}}$</td>
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<td>50 - 0.1</td>
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<td>-</td>
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<td>(NTU)</td>
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<td>Colour$_{\text{in-out}}$ (abs)</td>
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<td>Coliforms$_{\text{in-out}}$</td>
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<td>100%</td>
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<tr>
<td>(cfu 100ml$^{-1}$)</td>
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<td>100%</td>
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<tr>
<td>TDS$_{\text{in-out}}$ (ppm)</td>
<td>Chuk</td>
<td>765</td>
<td>1050 - 50</td>
<td>-</td>
<td>1180 567</td>
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<td>2200</td>
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<td>TSS$_{\text{in-out}}$ (ppm)</td>
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<td>&lt; 2</td>
<td>&lt; 1 - 0.05</td>
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<td>TOC$_{\text{in-out}}$ (ppm)</td>
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<td>pH$_{\text{in-out}}$</td>
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<td>Resistivity$_{\text{in-out}}$</td>
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<td>Hardness$_{\text{in-out}}$ (ppm)</td>
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<td>Alkalinity$_{\text{in-out}}$</td>
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<td>Silica$_{\text{in-out}}$ (ppm)</td>
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<td>SO$_{4}^{\text{in-out}}$ (ppm)</td>
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<td>Na$_{\text{in-out}}$ (mg Na$\text{L}^{-1}$)</td>
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<td>Ca$_{\text{in-out}}$ (mg Ca$\text{L}^{-1}$)</td>
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<td>103</td>
<td>30 - 0.5</td>
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<td>P$_{\text{in-out}}$ (mg P$\text{L}^{-1}$)</td>
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Calcium: $^2$ total, $^b$ calcium; chlorine: $^c$ total, $^d$ free; alkalinity: $^e$ total, $^f$ HCO$_3^{-}$; conductivity.
5.13 Reference material

These case studies have been researched by discussion with the end client and/or the supplier combined with information found on the internet or in published articles. The following list includes all sources except direct discussion as outlined in the Acknowledgements section. Web pages referenced were accessed on 12 November 2002.


Index

A
acetates 138, 139
acetic acid 69, 90, 112
acid baths 265, 266
acid cleaners 68, 69, 249, 273
acid dosing 64, 90, 239
acidified metabisulphite 70
acidity 188
acrylic fibres 143
acrylic monomers 21
acrylonitrile membranes 263, 272
activated carbon 67, 148, 149, 229, 245, 246, 266, 267
activated sludge (AS) processes 30, 44, 107, 119, 147, 148, 206, 244, 259
Adams-Millis Co. 149
adsorption 7, 45, 56, 67, 121, 122, 124, 141, 146–148, 267
aeration 30–31, 58, 67. see also activated sludge (AS) processes
aerosols 91
air conditioning 6, 76
air sparging 118
aldehydes 147
algorithms for pinch problem 201
alkaline cleaners 68, 69, 249, 273
alkalinity 5, 81, 217, 275
cooling and boiler feedwater 81, 86, 90, 94, 275
alkylsulphonates 69
Alpheus Environmental 228, 272
alum 90
alumina 30
aluminium 22, 60, 62, 81, 93, 115, 265
American Corporation 149
American Process 198
ammonia 81, 94, 96
ammonium 69, 70, 96
anaerobic treatment 146, 147, 258
Anglian Water Group (AWG) 228
anionic dyes 154
anionic trash 108
anisotropic membranes 16, 17, 19
Anopore filter 22
antistatic agents 137, 138, 139
AOX (adsorbable organic halogens) 107, 108, 109, 113, 114, 204
Apeldoorn, Netherlands 241–244, 270–275
apyrogenic water 5
Argo Analyser 90
arid regions 2, 5, 9
Arizona (Palo Verde nuclear power plant) 98
Aspen Equation Modeler 198
Aspen Properties 198
Aspen Technology 198, 199
Aspen Water (software) 198
ASTM, and fouling propensity 60, 61, 64, 65
ASTRASAND filters 259
asymmetric membranes 16, 19
Australia 160, 232–237, 270
Austria 6
automotive industry 4, 262–265, 269–275
B
backflushing 52, 56–57, 59, 66–67, 222–225
food and beverage industry 273
power industry 234, 273
pulp and paper industry 245
see also cleaning
backflushing water 160, 167
bacteria 17, 51, 65–66, 264, 266
automotive industry 264
pharmaceutical industry 165, 170
power industry 90–91, 229, 235
pulp and paper industry 115
bactericides 69–70
barium 61, 62, 64, 68, 175, 241
Bechtel 217
Belgium 6, 257–261
Bend Research 22
Borgen generating facility (New Jersey) 98
beverage industry 4, 76, 159–163
bicarbonate 69, 81, 241
Bigelow Carpets 149
biochemical oxygen demand see BOD

BOD (biological oxygen demand) 237
backflushing 52, 56–57, 59, 66–67, 222–225
food and beverage industry 273
power industry 234, 273
pulp and paper industry 245
automotive industry 264
pharmaceutical industry 165, 170
power industry 90–91, 229, 235
pulp and paper industry 115

biocides 66, 68, 69–70, 84, 91
biodegradability 138–139, 141, 146–148, 202, 250
biodegradability 65–66, 89, 90–93, 162–163, 167, 265
biological matter 58, 67, 68
in cooling and boiler feedwater 84, 85, 88, 90–93, 233
see also organic matter
biological treatment of food washing waste 255–257
depollution treatment 115, 116, 119, 248
of textile process effluents 140, 146–148, 150, 206, 251
black liquor 106, 116, 119
effluent treatment 115, 116, 120, 125, 241–244
bleaching of textiles 137–140, 145, 202, 204, 241–244
process modifications 148, 149
blinding 35, 96
blocking models 44–45
blood 17, 51, 52
blowdown water 85–88, 93, 94–95, 96–99, 100, 169
BOD
cooling and boiler feedwater 81, 95, 230–231, 235
food and beverage industry 255, 257, 275
pulp and paper industry 109, 114, 115, 275
textile industry 134, 135–140, 145, 146, 149, 150, 202–203, 204, 206
boiler feedwater 10, 33, 77, 80–81
food and beverage industry 159, 161, 162, 255
from municipal effluent 93, 228–241
pulp and paper industry 247
Borregaard sulphite pulp mill (Norway) 116
bottle washers 159–60
boundary layer 45, 48, 49, 50–51
Bounds editor 198–199
Brazil 102
brewing industry 4, 159, 161
brightness reversion 115
brine 239–240, 253, 266, 269
C
C30F filter 121, 122, 123
CAD see computer-aided design
cadmium 8, 62
cake layer 40–41, 43–51, 56
accumulation 35, 35–36, 56
mass transfer control 43–52
calcite 61
calcium 83, 86, 89, 94, 96, 112, 229, 241, 275
calcium carbonate scalants 51–52, 60, 61, 62, 64, 67, 68, 89, 96, 175, 181
calcium fluoride 62
calcium hydroxide 62
calcium phosphate scalants 62, 89, 231
calcium sulphate scalants 62, 64, 68, 86, 89
calcium sulphotannate 62
calcium carbonatescalant 51–52, 60, 61, 62, 64, 67, 68, 89, 96, 175, 181
calcium fluoride 62
calcium hydroxide 62
carbon dioxide acidifier 90
carbon dioxide permeation 61, 81, 160, 220, 242
carbon dioxide removal 96, 179, 220, 239
carbon filters see activated carbon
carbon fines 59, 229
carbonate scale 59, 60, 61, 64, 67, 68, 81, 86, 89, 90, 115, 175
carbonising 138, 140
carboxylates 21, 69, 122
carboxymethyl cellulose 135, 138
carriers 19
cartridge filters 10, 24–25, 33, 60, 67, 246, 266
cartridge microfilters, boiler feedwater 10. 33
cassettes 24, 30
catalytic oxidation 67
cautic soda 68, 69, 179
food and beverage industry 160
microelectronics industry 266
power industry 90, 234
pulp and paper industry 105, 246, 273
textile industry 136, 137, 140
cellophane production 8
cellulosic fibres 143
cellulosic matter 104, 140, 244, 246
cellulosic membranes 66, 120, 162, 248, 249, 272
cellulose acetate (CA) 19, 20, 22, 162, 175, 234, 240, 272
Ceramen 22
ceramic membranes 16, 17, 18, 19, 22–23, 30, 70, 154
ceramic tubes 28
CEVI 200
chelants/chelators 68, 69, 89, 90, 139
chemical dispersants 91–92
chemical finishing of textiles 141–143
chemical oxidation 20–21, 67, 137, 251, 253
chemical oxygen demand see COD
chemical pulping processes 102, 104, 105, 108, 114–115
chemical reduction 253
chemicals industry 6, 19, 76, 92–93, 209
chemothermomechanical pulping (CTMP) 104, 105, 109, 114
Chevron Oil Refinery (Oakland, CA) 98
China 102, 132, 147
Chirk, North Wales 244–248, 270–275
chloramine 162–163
chloride 81, 83, 94, 115, 170, 241
chlorinated compounds 139
chlorinated rubber 139
chlorination 67, 160, 162, 229, 259
chlorine 20, 22, 65, 66, 67, 69–70, 89, 91, 235, 266
- bleaching of pulp 104, 105, 106–107
- in mechanical pulp water 112
- residual 83–84, 162
chlorine dioxide 104, 105, 107, 253
chloramines 139
chloroether 21
chloroform 22
chromium 62, 134, 145, 146, 251–254
CIP (clean in place) 6, 28, 70, 160, 161, 167–168, 169, 230, 239, see also cleaning
circulation water see white water
citric acid 68, 69, 90, 235
clarification 206, 229, 238
clays 32, 51, 60, 122, 123, 251

- cleaning 28, 32, 52, 56–70
- automotive industry 262, 263–264, 273
- food and beverage industry 256, 259, 260, 273
- power industry 214, 215, 236, 273
- pulp and paper industry 117, 121, 246, 249, 250, 273
- textile industry 252–253, 273
see also backwashing; CIP
clogging 55, 59
coacluation 59, 67, 271

- food and beverage industry 162
- microelectronics industry 266
- power industry 96, 229, 238
- pulp and paper industry 125
- textile industry 146–148, 206
coarse bubble aeration 30–31, 58
coating materials 102, 105, 110, 115, 117–118, 120, 122–124, 150, see also dyeing industry: pigments
cobalt sulphide 62
Coca Cola Amatil plant (Australia) 160
COD 5, 81, 275
- cooling and boiler feedwater 81, 230–231
- food and beverage industry 258, 260, 275
- pulp and paper industry 107–109, 111–114, 116, 118–119, 243, 244, 247, 249, 275
Colflo 253
colliforms 91, 166, 233, 235, 275
colloidal material 9, 17, 44, 50, 51–52, 60, 67, 113, 249, see also dissolved and colloidal material
colour 107, 108, 112, 204, 275
- discharge consents 134, 141, 145, 250, 253, 275
see also coating materials: dyeing; pigments
composites membranes 16, 19
computer-aided design for RO 61, 172–186, 217–220, 222, see also software
concentration gradient 34, 36
concentration polarisation (CP) 36–39, 45–51, 53, 61
- classical model 41, 45–50
- coefficient 175
- modified model 50–51
- in software design 172, 173, 174, 175, 176, 180, 181, 213
concentration ratio 86–88, 93
condensate cooling 76, 161, 232, 237
conductivity 5
- food and beverage industry 231, 260
- microelectronics 267, 268
- pharmaceutical industry 165, 167, 170
- power industry 229
- pulp and paper industry 112, 241, 275
continuous denionisation (CDI) 32
continuous microfiltration (CMF) 51, 118
conversion 33
cooling 3, 6, 7, 76–79, 85–100, 159, 160, 204, 232, 255
copper 62, 81, 93
corrosion 80, 81, 85, 89, 90, 93, 95, 115, 188
Costpro software 174, 182
costs 4–5, 269–271, 274
- automotive industry 264
- backflushing and cleaning 67–68, 69, 234
- EDR/RO system for zero liquid discharge 241
- food and beverage industry 160, 161–162, 257, 258, 260–261
- membrane materials and manufacture 17–18, 22, 175
- membrane use 10–11
- microelectronics industry 265, 269
- pulp and paper industry 116, 117, 119, 125, 242–244, 247
- of purification 3, 235–237, 269
- recirculating water treatment 88, 116, 117
- software for costing 182, 198, 202
textile industry 148–149, 154, 203–207, 254
cotton 136–137, 138, 139, 140, 143, 146, 202, 241–244
CR (cross-rotational) filter 31–32, 116, 117, 118, 121, 248, 272
creosols 138
- critical flux 39–40, 51, 121, 249
- cross-flow operation 33, 36, 45, 51
- and fouling 59, 67, 121
- and polarisation 38–39
- cross-flow velocity 174, 176, 180, 214, 242, 249, 263, 273
- cross-rotational filter see CR filter
cryptosporidia 9, 17
current density 34
cycles of concentration 86–88
cyclohexanol 138
D
DAF see dissolved air flotation
Membranes for Industrial Wastewater Recovery and Re-use

dairy industry 4, 61, 76, 159, 161
DCS see dissolved and colloidal substances
dead-end operation 33, 36, 44–45
cleaning 70
design example 222–225
and fouling 56–57, 59
semi-dead-end mode 229, 259
dealkalisation 5, 160
debarking 104, 106, 107, 109
dehlorination see chlorine
deep bed/depth filtration 60, 67, 229, 238, 239, 245, 259, 266, 271
deonionisation 5, 9, 263, 265–269, see also electrodionisation
demineraliser feedwater 232–233, 239, 240, 270
Denmark 6, 200
dense membranes 15, 16, 52, 61, 119
depth filtration see deep bed/depth filtration
dersonisation 115, 125, 266, 267
Desal 22, 161, 174
desalination 9, 38, 54, 178, 220–222
Desalination Systems 153
design examples 213–225, see also computer-aided design
desizing 136, 138, 144, 146, 150, 202, 204, 207
detergents 67, 68
pharmaceutical industry 167
power industry 234
pulp and paper industry 249, 273
textile industry 136, 138, 140, 149, 150
dialysis 7, 9, 14, 39, see also electrodialysis
diffusion 36, 46, 47, 48, 51, 181
disc (mechanical) filtration 108–110, 114
disinfection 70, 83–84, 91, 233
dissociation 122, 172
dissolved air flotation (DAF) 67, 118, 244, 266
dissolved and colloidal substances (DCS) 108, 110, 112, 115, see also colloidal material
dissolved organic material 10, 17, 112
dissolved oxygen 94, 139
dissolved solids 81, 83, 95, 97, 112, 134, 140, 235, see also TDS
distillation 166
dolomite 62
domestic water 2, 3, 6
Donnan equilibrium model 41, 42, 43
Dora Creek wastewater treatment plant (Australia) 232
Doswell combined cycle power plant (USA) 237–241, 271–275
Dow 22, 121, 174
Dow Danmark Rina 22
drift water 85, 87
driving force 32, 34–349
“dumped” water 168
Duratherm Excel 161
dyeing industry 4, 139, 140–141, 145, 146–148, 149
case study 250–254
effluent 133–134, 141, 150–154, 202
in water pinch case study 202, 203, 204, 206, 208
E
EDTA (ethylene diamine tetracetic acid) 68, 69, 90
electro-oxidation 22
electrodionisation (EDI) 32, 99, 167, 169
electrodialysis (ED) 9, 14, 16, 19, 26–27, 34, 35, 43, 95, 125
and concentration polarisation 37–38, 49
design example 220–222
energy demand 57–58
modules 24, 25–27, 32
spacers 58
staging 52, 54
electrodialysis reversal (EDR) 58, 96, 238, 239–241
electrolysis 147, 253
electroneutrality 177
electrophoretic painting 3, 262–265
electrophoresis 177
energy demand 55–58, 67, 103, 176, 181, 187
in design examples 213–217, 220–222, 225
environmental awards 231, 248
environmental legislation and guidelines
pulp and paper industry 113–114
textile industry 133, 144–146, 154
see also legislation and guidelines
environmental preservation 265
enzymes 136, 138
enzymic cleaners 68, 69, 260
EPS (extracellular polymeric substances) 65, 66, 67
equilibrium chemical models 90
equilibrium thermodynamics 61–64, 173, 175
Eraring Power Station (Australia) 232–237, 270–275
Esmil Process Systems 247, 248, 272
estuarine water use in power industry 80–84
etching 18, 19, 22–23, 265
ethylchlorophosphates 143
ethylene diamine tetracetic acid (EDTA) 68, 69, 90
ethylene oxides 69, 139
Europe 6, 9
pharmaceutical industry 164, 165
pulp and paper industry 102, 108, 112, 117, 241–244
textile industry 132, 144–146
see also European Union
European Institute of Membranes 174
European Membrane Society 14
European Pharmacopoeia (EP) 164, 165, 166, 170
European Union
Integrated Pollution Prevention and Control (IPPC) Directive 113
potable water 144
and textile industry 132, 145 146
Urban Waste Water Treatment Directive (UWWTD) 9

evaporation 85, 87, 88, 93, 95, 160, 237–241
dairy industry 161
pulp and paper industry 103, 115, 117, 119
extracellular polymeric substances (EPS) 65, 66, 67
extracelluar processes 38, 39, 49
extrusion (stretching) 18, 19

F
Faraday equation 220

fats, oils and grease (FOG) 66, 138, 150, see also
oil and grease
fatty acids 108, 112, 121, 123, 249
feed 33, 53
feedwater
composition 174–175, 185, 217
pH 174, 175, 177
quality 177
temperature 174, 175, 177

Fenton's reagent 147

fibre, in pulping process 105, 108
fibre membranes 28–29
fibreboard (MDF) production 118–119, 244–248
fibrous materials 66
filamentous material 59
fillers 102, 122
film theory 41, 47, 173, 213
FilmTec 19, 121, 153, 174, 182, 185
finishing of textiles 141–143, 145, 146, 202
Finland 6, 112, 121, 248–250
fireproofing/flameproofing of textiles 139, 145
Flag Fen (UK) 98, 228–231, 270–275
flat sheet membranes 16–17, 121
flocculation 65, 118, 125, 146, 147, 238, 245, 251, 253, 259
Florida 98, 99
flooding
pulp and paper industry 103, 108, 114, 118, 242, 244
textile effluent 146, 150
flow rates
case studies 273
design examples 214–216
modelling 188–197
Fluid Systems see Koch
fluoride 170, 241, 267
fluorinated polymers 70
fluorocarbons 139
flow 32–33, 34, 36–37, 39–48, 51, 121, 122, 174, 176, 271
decline 44
and particle shape 123–124
range 27–71
in software design 174, 176, 178, 180–181
see also critical flux
foam balls 28
foaming 81
food industry 4, 6, 76, 159, 161, 209
case studies 254–261, 270–275
forest resources 102, 248
formaldehyde 68, 69–70, 139, 141
formate 118, 139
formic acid 112
Forsell et al. 250–254

G
gas separations 18, 19, 233
gas transfer (GT) 9, 14, 16, 19
gaseous water acidifiers 90
gel layer formation 35, 37, 48, 122, 172
gelatine 138, 139
Germany
automotive industry 262–265, 270–275
bottle washing wastewater recycling 161
paper mills 102
software production 174, 200
textile industry 132
water consumption 6

gluconic acid 69
glue residue 105
glycerol 138
grafting 20, 21
Grangemouth biofouling studies 92–93

Green Bay Packaging Inc (USA) 115
groundwater 80, 83, 242, 243, 257–258, 259, 261
Guadeloupe power plant (Texas) 99

H
Hagen-Poiseule equation 41, 56
Hamburg-Harburg Technical University 200
hardness 5, 38, 67, 81, 188, 217, 221
food and beverage industry 162, 260
pharmaceutical industry 167

Index 283
power industry 81, 275
pulp and paper industry 241, 247, 275
textile industry 207
heat exchange 6, 242, 244, 258, 269, 270
heavy metals 8, 165, 253–254
hemicelluloses 104, 108, 138
hemodialysis 7, 14
herbicides 17, 136
high-purity steam 76
high-purity water 5, 54, 166, 167, 170
case studies 228–237, 266–269
high-shear see shear enhanced processing
holding time index 87
hollow fibre modules 23, 28–29, 118, 150, 173, 222, 229, 233, 256, 259, 272
homogeneous membranes 16–17
humic acids 21–22, 260
Hungary 6
Hunter Sewage Project (Australia) 232
Hydronautics 174, 177, 182, 259
hydraulic resistance 40, 45, 52, 67, 172, 176, 215, see also membrane resistance
hydraulic staging 54, 221
hydrochloric acid 68, 69, 90, 137, 179, 219, 273
hydrofluoric acid 265
hydrogen peroxide 70, 251, 267
bleaching of pulp 104, 105, 108
bleaching of textiles 137, 139–140, 241
hydrogen sulphide 90
hydrolysis reactions 64, 233
hydrophilic membranes 22, 120, 249, 256, 239, 272
hydrophobic membranes 120, 272
hydrophobicity 20, 21, 112, 120, 121, 123–124, 272
Hydrotech 229
hydroxides 90
hypochlorite 68, 69, 137, 222, 230, 273. see also sodium hypochlorite

I
immersed membrane systems 58, 59, 70, 222, see also submerged membrane bioreactors
incineration 116, 117, 246
International Microelectronics (North San Jose, CA) 99
ion balance 175
ion depletion 35, 37–38
ion exchange 65, 90, 95, 96, 167, 231, 239, 266, 267
ion exchange dealkalisation 65, 160
ion exchange membranes 10, 16–17, 19
ion exchange resin 233
ion exchange resin beads 32
ionic product 61
ionics 22, 28, 32
lonpure 17, 32
ions 5, 15, 26
iron 58, 60, 62, 69, 81, 175, 260
compounds 62, 69, 115, 251, 233
cooling and boiler feedwater 81, 93, 94, 96
pulp effluent 115
irradiation 21. see also UV radiation
irrigation 2, 3, 6
Italy 6, 132, 144, 149
IUPAC 15

J
Japan 2, 254–257, 270–275
Japanese Pharmacopoeia (JP) 164
jute 138

K
Kanagawa Prefecture (Japan) 254–257, 270–275
kaolin 122
KBC Process Technology Ltd 187
Keram module 27
Kerasep ceramic membrane element 29
kidney dialysis 7
Kirkniemi (Finland) 121, 248–250, 270–275
Korea 102
Korony-Carman equation 44, 51
Kraft pulping process 104, 106–107, 109
Kronospan Ltd (UK) 244, 247
Kubota 11, 24, 30

L
laboratories 6, 8, 23
Lancashire medical textile company 149
landfill 115, 117, 238
Langeler Saturation Index (LSI) 5, 63, 64, 185, 219
latex 51, 110, 150
laundries 6, 59
lead sulphide 62
Legionella 90, 91
legislation and guidelines 2, 3, 271
cooling and boiler feedwater 93
food and beverage industry 258, 261
pharmaceutical industry 164–166
pulp and paper industry 113–115, 242
textile industry 133, 144–146, 134, 250, 253, 271
Lévêque solution 48–49, 50, 51
lignins 104, 105, 106, 108, 112, 123, 244, 246
lignosulphonates 116
lime/lime-soda softening 65, 67, 90, 95, 96, 251
limiting flow rate concept 188–197
LCv&que solution 48–49, 50, 51
Linnhoff March (LM) software see WaterTarget
lipophilic extractives 108, 112
Livingston (NEC Semiconductors (UK) Ltd) 265–269, 270, 271
LM (Linnhoff March) software see WaterTarget
Lockheed-Martin Electronics operation (Ocala, FL) 98
LSI see Langelier Saturation Index
lubricants 135, 137, 140
Luxembourg 6

M
M-Real Kirkniemi paper mill (Finland) 121, 248–250, 270–275
McKinley Paper plant (New Mexico) 118
Macroorb 148
magnesium 83, 86, 89, 90, 94, 96, 241
magnesium salts 61, 62, 64, 89, 90, 175
make-up water 85, 87, 94, 96–98, 238–239.
see also boiler feedwater
maltings steep water 161
manganese 62, 81, 93, 235
mass balance 33, 86, 173–174, 246
and critical flux 39
mass transfer control 41–52
mass transfer promotion 58, 173
mass transfer theory and modelling 40–41, 188
Matlab simulation tool 200
MBRs see membrane bioreactors
MDF (medium density fibreboard) production
meat industry 76
mechanical pulping processes 102, 104, 105, 106, 108–110, 109, 112, 114, 120
medical applications 6, 19, 149, 164, 166
melamine resins 139
membrane
definition 14
configurations 23–32
manufacture 17–23
mass transfer control 41–3
materials 15–23, 174
morphology 16
permeability 40, 56–57, 66, 67, 172
processes 7–11, 14
resistance 15, 32, 35
stability/integrity 20, 174, 175, 230, 234, 264
structure 14–17
membrane bioreactors (MBRs) 30–31, 44, 45, 46, 47, 52, 162, 242–243, see also submerged membrane bioreactors
Memcor 10, 118, 161, 233
mercerising 138, 140, 145, 149, 202, 204, 205
mercuric salts 62
metallic catalysts 140
metallic corrosion see corrosion
metallic membranes 16, 24
metallic scalant 67, 68
metals industries 6, 8, 79
Metso 272
Mexico 112
microbially induced corrosion (MIC) 90
microbiological safety 83–84, 167
microelectronics industry 6, 9–10, 79, 99, 265–269, 270, 271
microfiltration (MF) 9, 10, 11, 14, 17, 19, 20, 51, 60
beverage industry 161
boiler feedwater 10, 235
and concentration polarisation 37
continuous (CMF) 118
design example 222–225
membranes for 15, 16, 18, 19, 23–24
municipal water treatment 33, 229–230, 233–235
textile process 151–154, 206
microrotation 108
microorganisms 2, 5, 65–66, 90, 91, 166, 229, 249
microstrainers 59
mineral salts
cooling and boiler feedwater 81, 82, 83, 86, 88, 89–90, 95
formation 37, 38, 61–65, 107
rejection 19, 67, 239, 249
see also scalants
Mitsubishi Rayon Co Ltd 272
models, for mass transfer in membrane processes 40–41, 172, see also water pinch analysis
modified starches 135, 136, 138
MoDo Husum sulphate pulp mill (Sweden) 117
molecular weight cut-off (MWCO) 15, 117, 153
Monsanto Prism 173
mothproofing 139
municipal effluent 2–3, 10, 31, 33, 59
reclamation and reuse 93, 98, 228–237
treatment of industrial water 3–4
MWCO see molecular weight cut-off

N
N food plant (Japan) 254–257, 270–275
Nadir Filtration GmbH 122, 248, 272
nanofiltration (NF) 9, 14, 17, 20, 40, 43
for caustic recovery 160
ceramic membranes 18
and concentration polarisation 37
design software 174, 183
dyestuff removal 151–154
and fouling 60
make-up and blowdown water 97
maltings steep water 161
membranes for 15, 16, 19, 22, 27
potable water 21–22
pulp and paper industry 119, 121, 249
staging 52, 53
naphtol 139
Natal University Pollution Research Group 200
natural gas power station 237–241
natural organic matter (NOM) 21, 22, 52, 59, 60, 65, 66, 67, 122, see also organic matter
NEC Semiconductors (UK) Ltd (Livingston) 265–269
Nernst-Planck equation 42
Netherlands 6, 241–244
New Jersey (Bergen generating facility) 98
New Logic 31
New Mexico (McKinley Paper plant) 118
NF200 membrane 121
nickel salts 62
nitrate 96, 165, 241
nitric acid 69
nitrogen 96, 107, 109, 114, 204, 275
Nitto 22
NOM see natural organic matter
Norit X-flow 259, 272
North Carolina 149, 154
Norway 116, 150
Norway spruce (Picea abies) pulping 108
nuclear power stations 98, 99
Nuclepore membranes 22–23
nutrients 65, 81, 82, 83, 91, 95, 96, 107
nylon fibres 143
nylon-based filter 21

O
oil and grease 134, 136, 137, 140, 146, see also fats, oil and grease
Oklahoma University 200
once-through cooling systems 78, 79, 93, 95, 98
ophthalmic products 166
optimisation, pinch methodology 201, 206–209
organic acid cleaners 69
organic acid foulants 69, 90, 235
organic carbon see TOC
organic matter 60, 66, 67, 68, 81
food and beverage industry 161
microelectronics industry 266
power industry 81, 82, 91, 95, 97, 233, 256
pulp and paper industry 108, 121, 125
see also biological matter; natural organic matter (NOM)
organic nanofiltration membranes 22
organic polymeric media 24
organic solvents 22, 66, 150
organochlorine chemicals 137
organophosphates 137
ortho-chloromethylated polymer 21
orthophosphates 94
Osmonics 22, 119, 121, 133, 174, 179, 185, 246, 251, 272
osmotic pressure 37, 43, 54, 172, 173, 174, 175
oxalate 138
oxalic acid 69
oxidants 67, 68, 69, 104, 105
oxidation 20–21, 91, 93, 146–148, 251, 266, 267, see also chemical oxidation
ozonation 84, 91, 104, 125, 147, 148, 149, 206

P
PA 50H membrane 123
Pacific Power 232, 236
packaging 115, 159–160, 166
paint see coating materials; pigments
Pull 21, 31, 161, 272
Palo Verde nuclear power plant (AZ) 98
paper industry 3, 4, 105–106, 113, 241–244
industrial water demand 4
product types 102, 106, 108, 113, 117, 241
see also recovered paper
Paques 272
paraffin wax 139
parasite-control chemicals 137
particle bombardment 18, 22
particle count 5, 230
particle shape 123–124
particle size 51
particle transport 52, see also mass transfer
Pasfrost vegetable processing plant (Passchendaele, Belgium) 257–261
pasteurisers 160
pathogen removal 97
PCI Membranes 28, 272
PCI Memtech 160, 161
PECO Energy nuclear generating station (Pennsylvania) 99
pectins 66, 136
PEEK 22
Pennsylvania, PECO Energy nuclear generating station 99
pentachlorophenols 136, 138, 143
peptic matter 138
permanganate value (PV) 166
permeability 15, 19, 37, see also membrane permeability
permeate flux 174, 175, 177, 181, see also flux permeate quality 174, 181, 231
permeate stream 33
permeability 15, 16, 19, 20, 34, 177, 178
gel layer 37
peroxyacetic acid 70
pervaporation (PV) 9, 14, 16, 19
PES 50H membrane 123
pesticides 17, 137
PET 19, 22–23
Peterborough (UK) 228
petrochemical industry 76, 92–93, 98, 209
petroleum spirit 138
PH 5, 19–20, 22, 38, 174, 175, 273, 275
automotive industry 264
food and beverage industry 160, 162, 163, 261, 275
and fouling 58, 61, 63, 64, 69
power industry 81, 86, 88, 90, 94, 96, 229, 230, 239, 240, 241, 275
pulp and paper industry 106, 112, 122, 242, 275
in software design 174, 175, 179, 217
textile process effluent 134, 145, 146, 275
pharmaceutical industry 6, 9, 61, 163–170, 209
phase inversion process 19
phenols 122, 134, 139, 145, 146
phosphates 69, 81, 86, 89, 90, 94, 267
phosphonates 65
phosphonocarboxylic acid 69
phosphoric acid 69
Index 287

phosphorus 65, 90, 96, 107, 108, 109, 114, 204, 275
photicatalysis 147
pigments 3, 102, 105, 110, 262–265, 270. see also coating materials
pinch analysis 5, 186, 187, 199. see also water pinch
pinch point 189, 194
PLAIDE UF/MF module 25–26
plasma treatment 20, 21
plate-and-frame modules 23, 24–27, 245, 263, 272
plating plant 8
Poland 201
polishing 9–10, 24, 98, 149, 150, 161, 167, 247, 259, 266
pollution control 14, 93, 144–146, see also environmental legislation
polyacrylates 65, 150, 151
polyacrylonitrile (PAN) 20, 22
polyamide membrane materials 19, 69, 162, 230, 246, 256, 259, 272
aromatic 19, 20, 69, 175
polyamide textiles 139
polyaramide/polysulphone 249
polyaryletherketone (PAEK) 22
poly carbonate (PC) 19
poly(carboxylic acids) 65, 150, 151
polyelectrolytes 245
polyethylene membranes 256, 272
polyethylene terephthalate (PET) 19, 20, 22, 256
polygalacturonic acids 108
polyvinylpyrrolidone (PVP) 259, 272
poresize 15, 16, 18, 272
porous membranes 15, 16, 17, 19, 22, 24
Portugal 200
Posidyne nylon-based filter 21
potable water 143, 144, 166, 266
reserves 77
treatment 21, 166
potassium 83, 170, 241
power industry 4, 76–101
case studies 228–241, 270, 275
legislation and guidelines 3, 6, 93
see also boiler feedwater: cooling systems
PP (polypropylene) 20, 206, 233
pre-sterilisation 66
precipitation 146–148
precipitation 59
predictive models for mass transfer 40–41
pressing process 19
pressure decay tests 230, 234
pressure-driven processes 16, 38–39, 48, 49, 55–57, 60, 61
pressure-sealed modules 24–26
precipitation 52, 59–60, 65, 66, 67, 175, 185, 271
cooling and boiler feedwater 80, 90, 228–232, 238
food and beverage industry 162, 255
software design 175, 179–180, 219–220
textile processes 146, 202, 251
wood and pulp effluent 116, 118–119, 125, 245–246
printing, of textiles 135, 145, 202, 204, 206
printing ink 105, 111, 119
process design and operation 52–70, 176–178
process performance, definitions 32–34
process water quality requirements 6
proteins 17, 20, 44, 47, 66, 69
PTFE 19, 20
pulp and paper industry 3, 4, 102–131
case studies 241–250, 270–275
pulp and paper industry 77–84, 91, 93, 95–100, 228–237
microelectronics industry 265–269
power industry 77–84, 91, 93, 95–100, 228–237
textile industry 148, 149
see also recycling
recovered paper (RCF) 102, 104–105, 111, 113, 114

Q
quaternary ammonium compounds (QAC) 69, 70

R
Raytheon Corporation (St Petersburg, FL) 99
recirculating systems 78–79, 85–90, 93, 95, 98, 99
reclamation and reuse 2–4, 95, 269–271
microelectronics industry 265–269
power industry 77–84, 91, 93, 95–100, 228–237
textile industry 148, 149
see also recycling
recovered paper (RCF) 102, 104–105, 111, 113, 114
Membranes for Industrial Wastewater Recovery and Re-use

recovery 2, 31, 33, 53, 181, see also reclamation and reuse

recovery 2–3, 191, 194, 195

food and beverage industry 159–163, 254–261

microelectronics industry 265–269

pharmaceutical industry 163, 166–167, 168–170

plating plants 8

pulp and paper industry 114, 115, 119, 241–244, 248–250

textile industry 132–133, 146–154, 204–209, 250–254

see also closed-loop recycling; municipal effluent; reclamation and reuse

rejection 34

resin factory 8

resin removal 115, 125, 266, 267

resins

pulp and paper processes 108, 112, 115, 121, 123, 125, 244, 245, 246

textile processes 141, 145

resistance of interfacial region 35–36, 43–45, see also cake layer

resistance of membrane see membrane resistance

resistivity 5, 266, 269, 275

retentate stream 33, 53, 176

retention aids 108, 122, 125

reuse 2–3, 200, 269–271

of heat 241–244

see also municipal effluent; reclamation and reuse

reverse osmosis (RO) 9, 14, 15–17, 19, 20, 34

cleaning tanks 70

computer-aided design for plant 172–186 and concentration polarisation 37

design examples 213–220, 222

double-pass 166

energy demand 55–56

food and beverage industry 160, 161, 162, 256, 257, 259, 260

and fouling 55, 56, 66, 162

high temperature 161

leaky RO 16

make-up and blowdown water 97

membranes for 15–17, 19

pharmaceutical industry 165, 167, 169

pulp and paper industry 115, 118–119, 245–248

and scaling 60, 61, 64, 65

semiconductor industry 266, 268

software 61, 172–186, 217–220, 222

staging 52–54

substrates 20

textile effluent 132–133, 146–154, 204–209, 250–254

Reynolds number (Re) 48, 58, 214

Rhodia-Orelis 25–26, 29, 263, 272

risk analysis 2–3

municipal reuse 2–3

Rochem reverse osmosis module 24, 26

Rodata software 174

Rodesign software 174, 177, 185

ROP software 174, 175, 177, 178, 181, 218

Rupur 174

ROSA software 174, 182, 185

Rottneros Board Svaneholm (Sweden) 117

RWTH Aachen 174

Ryznar Stability Index (RSI) 63–64

Rzeszów University of Technology (Poland) 201

Saarland University (Germany) 174

saturation pH 63

scalants and scaling indices 35, 37–38, 58, 61–65, 115, 174, 175, see also specific scalants

scale formation propensity 61–64, 90, 173, 175, 176, 181

scale prevention 62, 64–65, 181, 239

in cooling and boiler feedwater 80, 82, 85, 86, 87, 88, 89–90, 95, 96, 233

Schmidt number (Sc) 48, 214

scouring 136–137, 138, 140, 146, 150, 202

screens 59, 229, 233, 251, 258

SDI (Silt Density Index) 5, 6, 60, 229, 233

seawater 80, 82, 83, desalination 9, 54, 178

SelRO 22

semiconductor industry 6.9–10, 265–269, 270, 271

sensitivity analysis 193

Severn Trent Water 146, 250

sewage treatment 2, 3–4, see also municipal effluent

shear rate 49, 51, 121

shear-enhanced processing 31–32, 58, 119–120, 121, 125

shear-induced diffusion 50, 51–52

Sherwood number (Sh) 48, 213–114

sieving 15, 40

silica 5, 17, 52, 59, 65, 67, 68, 69, 162, 275

cooling and boiler feedwater 81, 86, 89, 94, 96, 229, 247, 275

pulp effluent 115

silicon carbide substrate 30

silicon waste recycling 265

silicone resins 139

silk 138, 149

silt 59

Silt Density Index (SDI) 5, 6, 60, 229, 233

silver salts 62

Singapore oil refinery 98

singeing 135, 202

single pass with recycle 176

sintering 18, 19

sizes 105, 135, 136, 137, 150, see also desizing

soap industry 4

soaps 69, 136, 140, 148–149

sodium 83, 170, 229, 241, 275

sodium bisulphite 137

sodium carbonate 136, 140, 149
sodium chloride 172, 204, 232, see also desalination
sodium hexametaphosphate (Calgon) 64–65
sodium hydroxide 106, 253, 273
sodium hypochlorite 69, 137, 233, 273, see also hypochlorite
sodium hyposulphate 68
sodium sulphide 106
sodium thiosulphate 65, 104
softening 5, 60, 65, 67, 96, 167, 254, 261, see also lime
software

designs for RO 61, 172–186, 217–220, 222
sol-gel processes 18, 19
solubility 61, 62, 90, 173, 175, 188
solution-diffusion model 40–41, 42–43, 172
solvents 22, 66, 139, 150
Somicot 22
sonic tests 234
sorption-capillary flow 41
South Africa 147, 150, 154, 200
South Wigston (UK) 250–254, 270–275
spacers 58, 173
Spin 6
SPEEK see sulphonated poly(ether ether ketone)
SPEES see sulphonated poly(ether ether sulphone)
spiral-wound membranes 27–28, 60
food and beverage industry 256, 259
power industry 230, 234, 272
pulp and paper industry 119, 121, 249, 272
textile industry 252, 272
staging 52–54, 176, 178, 221
stagnant region 46–47, 176, 178, 221
starches 66, 122, 135, 136, 138, 146, 148, see also sizes
statutory requirements, 3, see also legislation
steam raising 6, see also boiler feedwater
steam sterilisation 70
steam in textile manufacture 204
stearates 139
steel 93, 151
sterilisation 19, 66, 259, 268
sterols 123
steryl esters 123
sticky residues in recovered paper 105, 113
Stiff and Davis Index 64
Stokes-Einstein equation 48
Stora Kabel 117
Stora Enso Uetersen 117
streptococci 233
stretching (extrusion) 18, 19
strontium 64, 175
submerged membrane bioreactors (SMBR) 11, 24, 30–31, 46, 58, 59, 150, 255–256, see also immersed membrane systems
substrates 20, 30
sugars 17, 108, 112, 116, 121, 249
stilt 188
sulphate 64, 65, 67, 83, 175, 275
cooling and boiler feedwater 81, 86, 89, 90, 94, 241, 247, 275
pulp effluent 113, 275
semiconductor industry 267
sulphate pulping process 104, 105, 117
sulphides 134, 145, 146, 175
sulphite pulping process 104, 105, 109, 115, 116, 125
sulphonated poly(ether ether ketone) see SPEEK
22
sulphonated poly(ether ether sulphone) see SPEES 22
sulphur dioxide acidifier 90
sulphuric acid 69, 90, 105, 137, 140, 179, 233, 251, 267
supported liquid membranes (SLMs) 18–19
surfactants 68, 69, 136, 137, 150, 242, 273
suspended solids 60, 67
food and beverage industry 256–257, 275
microelectronics industry 266
power industry 229, 233, 249, 275
pulp and paper industry 110, 112, 114, 117, 244, 275
textile industry 150, 202, 204, 206, 253
see also TSS
Swansea (University of Wales) 174
Sweden 6, 117
synthetic textiles 135, 138, 140

T
Taio Paper Co. 116
talc 123–124
TAMI 27
Tanapa waste-to-energy facility (Florida) 98
tartrate 139
TDS 5, 175, 217, 219–220, 222
food and beverage industry 160, 162, 275
power industry 231, 233, 240–241, 275
pulp and paper industry 275
textile industry 134, 253, 275
see also dissolved solids
Tellon Typer (Tetrace) membrane 123–124
temperature 87, 94, 204, 273
Texas (Guadelope power plant) 99
Texas Utilities (TXU) 228
textile industry 132–158
dye wastewater recycling case study 250–254, 270–275
water pinch case study 201–209
TFC see thin-film composite (TFC) layer
thermomechanical pulping (TMP) 104, 108, 109
thin-film composite (TFC) layer 16, 19, 69
thiosulphate 65, 104
threshold inhibitors 67, 173
titanates 139
titania 10
titanium 18
titanium dioxide microfiltration membrane 153, 154
Membranes for Industrial Wastewater Recovery and Re-use

TOC 5.175.183
microelectronics industry 266.267
pharmaceutical industry 165.166.170
power industry 94.229.241.275
textile industry 134.148
Toray 174.256
toxics 95.136.137.140.143.165.170
track-etched polymers 19.22-23
transmembrane pressure (TMP) 34.36.41.42.48.173-174.176.215
automotive plant 263.273
food industry 273
power industry 234.256.273
pulp and paper industry 242.246.249.273
TSS 5.275
cooling and boiler feedwater 81.95
pulp and paper industry 109.113.114.275
reclaimed water 81.95.230-231
textile processes 134.135.136.145.146.204.275
see also suspended solids
tubular modules 23.28.115-116.118.121.125.242.272
turbidity 5.81.235.275
turbulence, and biofilm thickness 65
turbulence promotion 23.24.28.30-32.37
U
UK 4-5.6.9
automotive industry 264-265
Flag Fen high-purity water production plant 98.228-231
food and beverage industry 161.162
Kronospan Ltd 244
pharmaceutical industry 168.169
pulp and paper industry 111-112.244-248
RO software 174
semiconductor industry 265-269.270.271
textile industry 132.146.150.250-254
ultrafiltration (UF) 9.14.17.19.20.33.60.67
automotive industry 262-265
and concentration polarisation 37
food and beverage industry 161.162.259.260.261
make-up and blowdown water 97
membranes for 15.16.17.18.19.22.24
pharmaceutical industry 166
in pulp and paper industry 115-118.125.248-250
semiconductor industry 266
textile process 150-154.204-205.206
and VSEP filter 121
ultrapure water 5.6.9-10.266.269
UMIST (Department of Process Integration) 187.199-200
see also Water (UMIST software)
urea 139
USA 2.3.229
pharmaceutical industry 164.166.168
Pharmacobio (USP) 164.166
power industry 91.93.98.99.237-241
pulse and paper industry 102.104.115.118
textile industry 132.133.144-145.149.154
UV (ultraviolet) processes 66.67.84.160.251.266.267-268
V
Valmet-Flotech CR filter 31-32.116-117
vanillin 116
van't Hoff equation 37.43.172.173
vapour compression evaporation 237-241
VHP security paper mill (Netherlands) 241-244
Vibrating Membrane Filter (VMF) 31.32.161
Vibratory Shear Enhanced Processing see VSEP
vinyl 21
viruses 17.91.233.235
viscose 138.139.143
Visual Basic-based software 183-184.200
Vito 184
Vivendi 10.272
Vivendi Ionpure 32
Vivendi Memcor 118.161
VMF (Vibrating Membrane Filter) 31.32.161
vortex flow 32.58
VSEP (Vibratory Shear Enhanced Processing) 31.32.119-120.121.123.124
W
Wado* software 200
Water (UMIST software) 187.193.194.195.198.199
water auditing 8.148.232
water balance 173-174.187
Water Close* (software) 198
Water Factory 21 plant (Orange County, CA) 3.229
water for injection (WFI) 9.165-170
water management 5-7.8
water pinch analysis 5.148.186-212
water quality 6-7.80-84.95.164-167
see also legislation and guidelines
water saving 192-195
water shortages 77
water splitting 38
WaterDesign software 200
waterprooiling 139.145
WaterTarget* software 187.191-199.201
waxes 135.136.137.138
white liquor 106
white water (circulation water) 108-110.114.116.117.120.248.249
Wigan dyehouse (UK) 149
Wigston sewage treatment works 146
WinCarol software 174
WinFlows software 174.179.185
wood 104, 122
wood plant effluent 118-119
wool 117, 118, 119, 120, 123, 126, 150

Y
Yokosuka (Japan) 254, 257

Z
Zenon 30
zeolite 148
zero liquid discharge (ZLD) 3, 8, 76-77, 95, 118, 271
  case studies 237-241, 244-248, 265, 269, 271
zinc 62, 253
zirconium 18, 30, 151
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