

FRESENIUS MEDICAL CARE CORP

Form 20-F

March 02, 2004

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UNITED STATES
SECURITIES AND EXCHANGE COMMISSION
Washington, D.C. 20549

FORM 20-F

REGISTRATION STATEMENT PURSUANT TO SECTION 12(b) OR (g) OF THE SECURITIES EXCHANGE ACT OF 1934

OR

ANNUAL REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

For the fiscal year ended December 31, 2003

OR

TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

Commission file number _____

FRESENIUS MEDICAL CARE
AKTIENGESELLSCHAFT

(Exact name of Registrant as specified in its charter)

FRESENIUS MEDICAL CARE CORPORATION

(Translation of Registrant's name into English)

Germany

(Jurisdiction of incorporation or organization)

Else-Kröner Strasse 1, 61352 Bad Homburg, Germany

(Address of principal executive offices)

Securities registered or to be registered pursuant to Section 12(b) of the Act:

Title of each class	Name of each exchange on which registered
American Depositary Shares representing Preference Shares	New York Stock Exchange
Preference Shares, no par value	New York Stock Exchange ⁽¹⁾
American Depositary Shares representing Ordinary Shares	New York Stock Exchange
Ordinary Shares, no par value	New York Stock Exchange ⁽¹⁾

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Securities registered or to be registered pursuant to Section 12(g) of the Act: None

Securities for which there is a reporting obligation pursuant to Section 15(d) of the Act: **7 7/8% USD Trust Preferred Securities due 2008, 7 3/8% DM Trust Preferred Securities due 2008, 7 7/8% USD Trust Preferred Securities due 2011, 7 3/8% Euro Trust Preferred Securities due 2011 and related guarantees**

(1) Not for trading, but only in connection with the registration of American Depositary Shares representing such shares.

Indicate the number of outstanding shares of each of the issuer's classes of capital or common stock as of the close of the period covered by the annual report:

Preference Shares, no par value 26,213,919

Ordinary Shares, no par value 70,000,000

Indicate by check mark whether the registrant (1) has filed all reports required to be filed by Section 13 or 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period that the registrant was required to file such reports) and (2) has been subject to such filing requirements for the past 90 days.

Yes No

Indicate by check mark which financial statement item the registrant has elected to follow.

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INTRODUCTION

Forward Looking Statements

This report contains forward-looking statements within the meaning of section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended, which are based upon our current expectations, assumptions, estimates and projections about us and our industry that address, among other things:

our business development, operating development and financial condition;

our expectations of growth in the patient population regarding renal dialysis products and services;

our expansion and acquisition plans and our capital expenditures budget;

the impact of our expansion on our revenue potential, cost basis and margins;

our ability to remain competitive in the markets for our products and services;

the effects of regulatory developments, legal proceedings and our settlement of government investigations into our business;

possible changes in government reimbursement policies and those of private payors;

our ability to develop and maintain additional sources of financing; and

other statements of our expectations, beliefs, future plans and strategies, anticipated development and other matters that are not historical facts.

When used in this report, the words *expects*, *anticipates*, *intends*, *plans*, *believes*, *seeks*, *estimates* and similar expressions are generally intended to identify forward looking statements. Although we believe that the expectations reflected in such forward-looking statements are reasonable, forward-looking statements are inherently subject to risks and uncertainties, many of which cannot be predicted with accuracy and some of which might not even be anticipated. Future events and actual results, financial and otherwise, could differ materially from those set forth in or contemplated by the forward-looking statements contained elsewhere in this report. Important factors that could contribute to such differences are noted in this report under *Business Overview* in *Item 4. Information on the Company*, *Item 5. Operating and Financial Review and Prospects* and *Item 8.A.7. Legal Proceedings*.

This report contains patient and other statistical data related to end-stage renal disease and treatment modalities, including estimates regarding the size of the patient population and growth in that population. These data have been included in reports published by organizations such as the Center for Medicare and Medicaid Services of the U.S. Department of Health and Human Services, the Japanese Society for Dialysis Therapy and the German registry Quasi-Niere. While we believe these surveys and statistical publications to be reliable, we have not independently verified the data or any assumptions on which the estimates they contain are based.

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Not applicable

Item 2. Other Statistics and Expected Timetable

Not applicable

Item 3. Key Information**Selected Financial Data**

The following table summarizes the consolidated financial information for our business for each of the years 1999 through 2003. We derived the selected financial information from our consolidated financial statements. We prepared our financial statements in accordance with accounting principles generally accepted in the United States of America and KPMG Deutsche Treuhand-Gesellschaft Aktiengesellschaft Wirtschaftsprüfungsgesellschaft, independent accountants, audited these financial statements. You should read this information together with our consolidated financial statements and the notes to those statements appearing elsewhere in this document and the information under Item 5. Operating and Financial Review and Prospects .

	2003 ^(A)	2002 ^(A)	2001 ^(B)	2000	1999
	(In millions)				
Statement of Operations Data:					
Net revenues	\$ 5,528	\$ 5,084	\$ 4,859	\$ 4,201	\$ 3,840
Cost of revenues	3,699	3,428	3,220	2,734	2,463
Gross profit	1,829	1,656	1,639	1,467	1,377
Selling, general and administrative	1,022	914	966	814	785
Research and development	50	47	36	32	32
Special charge			258		601
Operating income (loss)	757	695	379	621	(41)
Interest expense, net	211	226	223	216	218
Income (loss) before income taxes	546	469	156	405	(259)
Net income (loss)	\$ 331	\$ 290	\$ 63	\$ 212	\$ (249)
Weighted average of:					
Preference shares outstanding	26,191,011	26,185,178	26,035,330	19,002,118	9,023,341
Ordinary shares outstanding	70,000,000	70,000,000	70,000,000	70,000,000	70,000,000
Basic income (loss) per Ordinary share	\$ 3.42	\$ 3.00	\$ 0.65	\$ 2.37	\$ (3.15)
Fully diluted income (loss) per Ordinary share	3.42	3.00	0.64	2.36	(3.15)
Basic income (loss) per Preference share	3.49	3.06	0.70	2.43	(3.15)
	3.49	3.06	0.69	2.42	(3.15)

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Fully diluted income (loss) per Preference share					
Basic and fully diluted net income (loss) per Ordinary ADS	1.14	1.00	0.22	0.79	(1.05)
Basic and fully diluted net income (loss) per Preference ADS	1.16	1.02	0.23	0.81	(1.05)
Dividends declared per Ordinary share (⁽⁹⁾)	1.02(b)	0.94	0.85	0.78	0.69
Dividends declared per Preference share (⁽⁹⁾)	1.08(b)	1.00	0.91	0.84	0.75
Dividends declared per Ordinary share (\$) ^(a)		1.10	0.78	0.72	0.64
Dividends declared per Preference share (\$) ^(a)		1.17	0.84	0.78	0.69

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	2003 ^(A)	2002 ^(A)	2001 ^(B)	2000	1999
	(In millions)				
Balance Sheet Data					
Working capital	\$ 794	\$ 526	\$ 402	\$ 191	\$ (229)
Total assets	7,503	6,780	6,516	5,979	5,752
Total long-term debt (c)	2,354	2,234	2,165	1,611	1,618
Shareholders' equity (net assets)	3,244	2,807	2,617	2,679	2,002
Capital Stock	Preference shares Nominal				
Value	70	70	70	64	28
Capital Stock	Ordinary shares Nominal Value				
Value	229	229	229	229	229

(A) Includes the effect of an accounting change in 2002 relating to the adoption of SFAS No. 142, *Goodwill and Other Intangible Assets*, as of January 1, 2002 (see Note 1 of the Notes to our Consolidated Financial Statements).

(B) Includes the special charge to address 1996 merger related legal matters, estimated liabilities and legal expenses arising in connection with the W.R. Grace Chapter 11 proceedings and the cost of resolving pending litigation and other disputes with certain commercial insurers. You can find a more detailed discussion of this special charge in Note 3 of the Notes to our Consolidated Financial Statements.

(a) Amounts shown for each year from 1999 to 2002 represent dividends paid with respect to such year. The actual declaration and payment of the dividend was made in the following year, after approval of the dividend at our general meeting.

(b) Our managing board and our supervisory board have proposed dividends for 2003 of \$1.08 per Preference share and \$1.02 per Ordinary share. These dividends are subject to approval by our shareholders at our annual general meeting to be held on May 27, 2004.

(c) Total long-term debt represents long-term debt and capital lease obligations, less current portions and (i) at December 31, 1999 and 2000, the mandatorily redeemable preferred securities of Fresenius Medical Care Capital Trust, Fresenius Medical Care Capital Trust II, and Fresenius Medical Care Capital Trust III, (ii) at December 31, 2001, the mandatorily redeemable preferred securities of Fresenius Medical Care Capital Trust, Fresenius Medical Care Capital Trust II, Fresenius Medical Care Capital Trust III, Fresenius Medical Care Capital Trust IV, and Fresenius Medical Care Capital Trust V, (iii) at December 31, 2002 and 2003 the mandatorily redeemable preferred securities of Fresenius Medical Care Capital Trust II, Fresenius Medical Care Capital Trust III, Fresenius Medical Care Capital Trust IV, and Fresenius Medical Care Capital Trust V. On February 14, 2002, we redeemed the entire \$360 million aggregate liquidation amount of the trust preferred securities of Fresenius Medical Care Capital Trust.

RISK FACTORS***Risks Relating to Litigation and Regulatory Matters in the U.S.***

If we do not comply with the many governmental regulations applicable to our business or with the corporate integrity agreement between us and the U.S. government, we could be excluded from government health care reimbursement programs or our authority to conduct business could be terminated, either of which would result in a material decrease in our revenue

Our operations in both our provider business and our products business are subject to extensive governmental regulation in virtually every country in which we operate. The applicable regulations, which differ from country to country, relate in general to the safety and efficacy of medical products and supplies, the operation of manufacturing facilities, laboratories and dialysis clinics, the rate of, and accurate reporting and billing for, government and third-party reimbursement, and compensation of medical directors and other financial arrangements with physicians and other referral sources. We are also subject to other laws of general applicability, including antitrust laws.

Fresenius Medical Care Holdings Inc. (FMCH), our North American subsidiary, is party to a corporate integrity agreement with the U.S. government. This agreement requires that Fresenius Medical Care Holdings staff and maintain a comprehensive compliance program, including a written code of conduct, training programs, regulatory compliance policies and procedures, annual audits and periodic reporting to the government. The corporate integrity agreement permits the U.S. government to exclude Fresenius Medical Care Holdings and its subsidiaries from participation in U.S. federal health care programs if there is a material breach of the agreement that Fresenius Medical Care Holdings does not cure within thirty days after Fresenius Medical Care Holdings receives written notice of the breach. We derive approximately 43% of our consolidated revenue from U.S. federal health care benefit programs. Consequently, if Fresenius Medical Care Holdings commits a material breach of the corporate integrity agreement that results in the exclusion of Fresenius Medical Care Holdings or its

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subsidiaries from continued participation in those programs it would significantly decrease our revenue and have a material adverse effect on our business, financial condition and results of operations.

While we rely upon our management structure, regulatory and legal resources, and the effective operation of our compliance program to direct, manage and monitor these activities, if employees, deliberately or inadvertently, failed to adhere to these regulations then our authority to conduct business could be terminated or our operations could be significantly curtailed. Any such terminations or reductions could materially reduce our revenues with a resulting adverse impact on our business, financial condition and results of operations.

A reduction in U.S. government reimbursement for dialysis care could materially decrease our revenues

For the twelve months ended December 31, 2003 approximately 43% of our consolidated revenues resulted from Medicare and Medicaid reimbursement. Legislative changes may affect the reimbursement rates for the services we provide, as well as the scope of Medicare and Medicaid coverage. A decrease in Medicare or Medicaid reimbursement rates or covered services could have a material adverse effect on our business, financial condition and results of operations.

A change in reimbursement for or utilization of EPO could materially reduce our revenue and operating profit

Reimbursement and revenue from the administration of erythropoietin, or EPO, accounted for approximately 23% of dialysis care revenue in our North America segment for the twelve months ended December 31, 2003. EPO is produced by a single source manufacturer, Amgen Inc. Our current contract with Amgen covers the period from January 1, 2004 to January 31, 2006. A reduction in reimbursement for EPO, a significant change in utilization of EPO, a reduction of the current overfill amount in EPO vials, an interruption of supply or our inability to obtain satisfactory purchase terms for EPO after our current contract expires could reduce our revenues from, or increase our costs in connection with the administration of the EPO, which could materially adversely affect our business, financial condition and results of operations.

Creditors of W.R. Grace & Co. Conn. have asserted claims against us

We were formed in 1996 as a result of a series of transactions with W.R. Grace & Co that we refer to as the merger. At the time of the merger, W.R. Grace & Co.-Conn. had, and continues to have, significant liabilities arising out of product-liability related litigation (including asbestos), pre-merger tax claims and other claims unrelated to its dialysis business; in connection with the merger, W.R. Grace & Co.-Conn. and other Grace entities agreed to indemnify Fresenius Medical Care and its subsidiaries against all liabilities of W.R. Grace & Co., whether relating to events occurring before or after the merger, other than liabilities arising from or relating to National Medical Care's operations.

In addition, the merger was consummated as a tax free organization. Pre-merger tax claims or other tax claims that would arise if events were to violate the tax-free nature of the merger could ultimately be the obligation of our subsidiary, FMCH. In particular, W.R. Grace & Co. has publicly disclosed that its tax returns for the years 1993 through 1996 are under audit by the U.S. Internal Revenue Service, that it has paid tax and interest with respect to certain deductions taken prior to 1993 and that similar deductions were taken in the years under audit. Subject to certain representations made by W.R. Grace & Co.-Conn, FMCH and Fresenius AG, W.R. Grace & Co.-Conn. and other Grace entities also agreed to indemnify us against any such tax liability. W.R. Grace & Co.-Conn. and certain of its subsidiaries have filed for reorganization under Chapter 11 of the U.S. Bankruptcy Code (Grace Chapter 11 Proceedings). If W.R. Grace & Co.-Conn.'s (and its affiliates in and out of bankruptcy) obligations to indemnify us are terminated or limited as a result of bankruptcy proceedings, and if we are held liable for pre-merger obligations of W. R. Grace & Co.-Conn. or if the merger is determined to be a taxable transaction, our business, financial condition and results of operations could be adversely affected.

In 2002, claims were brought against W.R. Grace & Co. and FMCH by plaintiffs claiming to be creditors of W.R. Grace & Co.-Conn. (asbestos creditors), principally alleging that the merger was a fraudulent conveyance, violated the uniform fraudulent transfer act, and constituted a conspiracy. We are also engaged in litigation with Sealed Air Corporation (with which W. R. Grace & Co.-Conn. conducted a multistep transaction

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subsequent to the merger) over our entitlement to indemnification from Sealed Air for losses and expenses we incur relating to pre-merger tax liabilities of W.R. Grace & Co. and merger-related claims.

In 2003, the Company reached an agreement with the asbestos creditors' committees and W.R. Grace & Co. in the Grace Chapter 11 Proceedings to settle these fraudulent conveyance and tax claims. The settlement agreement has been approved by the U.S. District Court. The proposed settlement is subject to confirmation of a final plan of reorganization of W.R. Grace & Co. that meets the requirements of the settlement agreement or is otherwise satisfactory to us. If the proposed settlement with the asbestos creditors' committees and W.R. Grace & Co. is not confirmed in such a final plan of reorganization, the claims could be reinstated. If the claims are reinstated and the merger is determined to be a fraudulent transfer and if material damages are proved by the plaintiffs and we are not able to collect, in whole or in part, on the indemnity from any of our indemnitors, a judgment could have a material adverse effect on our business, financial condition and results of operations. We recorded a pre-tax accrual of \$172 million at December 31, 2001 to reflect our estimated exposure for liabilities and expenses related to the Grace Chapter 11 Proceedings. See Note 3 to our consolidated financial statements. For additional information concerning the Grace Chapter 11 Proceedings and the settlement agreement see Item 8.A.7 Legal Proceedings.

As health maintenance organizations and other managed care plans grow, amounts paid for our services and products by non-governmental payors could decrease

We obtain a significant portion of our revenues from reimbursement provided by non-governmental third-party payors. Although non-governmental payors generally pay at higher reimbursement rates than governmental payors, managed care plans generally negotiate lower reimbursement rates than indemnity insurance plans. Some managed care plans and indemnity plans also utilize a capitated fee structure or limit reimbursement for ancillary services.

As managed care programs have increased market share, there has been increased pressure to reduce the amounts paid for our services and products. These trends may be accelerated if future changes to the U.S. Medicare ESRD program require private payors to assume a greater percentage of the total cost of care given to dialysis patients over the term of their illness, or if managed care plans otherwise significantly increase their enrollment of renal patients.

If managed care plans reduce reimbursements, our revenues could decrease, and our financial condition and results of operations could be materially adversely affected.

Proposals for health care reform could decrease our revenues

Proposals to modify the current health care system in the U.S. to improve access to health care and control its costs are continually being considered by the federal and certain state governments. See Regulatory and Legal Matters-Reimbursement- U.S. for a discussion of the recently enacted Medicare Prescription Drug Modernization and Improvement Act of 2003. We anticipate that the U.S. Congress and state legislatures will continue to review and assess alternative health care reforms, and we cannot predict whether these reform proposals will be adopted, when they may be adopted or what impact they may have on us. Any spending decreases or other significant changes in the Medicare program could reduce our revenues and profitability and have a material adverse effect on our business, financial condition and results of operations.

Other countries, especially those in Western Europe, have also considered health care reform proposals and could materially alter their government-sponsored health care programs by reducing reimbursement payments. Any reduction could affect the pricing of our products and the profitability of our services, especially as we expand our international business. This potential development could have a material adverse effect on our business, financial condition and results of operations.

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Risks Relating to our Business

Our competitors could develop superior technology or impact our product sales

We face numerous competitors in both our dialysis services business and our dialysis products business, some of which may possess substantial financial, marketing or research and development resources. Competition could materially adversely affect the future pricing and sale of our products and services. In particular, technological innovation has historically been a significant competitive factor in the dialysis products business. The introduction of new products by competitors could render one or more of our products obsolete.

We are engaged in both manufacturing dialysis products and providing dialysis services. We compete in the dialysis services business with many customers of our products business. As a result, independent dialysis clinics, those operated by other chains and dialysis centers acquired by other products manufacturers may elect to limit or terminate their purchases of our dialysis products so as to avoid purchasing products manufactured by a competitor. In addition, as consolidation in the dialysis services business continues and other vertically integrated dialysis companies expand, the external market for our dialysis products could be reduced. Possible purchase reductions could decrease our product revenues, with a material adverse effect on our business, financial condition and results of operations.

We also compete with other dialysis products and services companies in seeking selected acquisitions. If we are not able to continue to effect acquisitions in the provider business in our International segment upon reasonable terms there could be an adverse impact on the growth of our business and our future growth prospects.

We face products liability and other claims which could result in significant liability

Health care companies are subject to claims alleging negligence, products liability, breach of warranty, malpractice and other legal theories that may involve large claims and significant defense costs whether or not liability is ultimately imposed. Health care products may also be subject to recalls. Although product liability claims and recalls have not had a material adverse effect on our businesses in the past, we cannot assure that we will not suffer one or more significant claims or product recalls in the future. Product liability claims or recalls could result in judgments against us or significant compliance costs, which could materially adversely affect our business, financial condition and results of operations.

While we have been able to obtain liability insurance in the past, it is possible that such insurance may not be available in the future either on acceptable terms or at all. A successful claim in excess of the limits of our insurance coverage could have a material adverse effect on our results of operations and financial condition. Liability claims, regardless of their merit or eventual outcome, also may have a material adverse effect on our business and reputation, which could in turn reduce our revenues and profitability.

If physicians and other referral sources cease referring patients to our dialysis clinics or cease purchasing our dialysis products, our revenues would decrease

Our dialysis services business is dependent upon patients choosing our clinics as the location for their treatments. Patients may select a clinic based, in whole or in part, on the recommendation of their physician. We believe that physicians and other clinicians typically consider a number of factors when recommending a particular dialysis facility to an ESRD patient, including, but not limited to, the quality of care at a clinic, the competency of a clinic's staff, convenient scheduling, and a clinic's location and physical condition. Physicians may change their facility recommendations at any time, which may result in the movement of our existing patients to competing clinics, including clinics established by the physicians themselves. At most of our clinics, a relatively small number of physicians account for the referral of all or a significant portion of the patient base. If a significant number of physicians ceased referring their patients to our clinics, this could reduce our dialysis care revenue and materially adversely affect our overall operations. Our operations are also affected by referrals from hospitals, managed care plans and other sources.

The decision to purchase our dialysis products and other services or competing dialysis products and other services will be made in some instances by medical directors and other referring physicians at our dialysis clinics and by the managing medical personnel and referring physicians at other dialysis clinics, subject to applicable

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regulatory requirements. A decline in physician recommendations or purchases of our products or ancillary services could reduce our dialysis product and other services revenue, and materially adversely affect our business, financial condition and results of operations.

If we are unable to attract and retain skilled medical, technical and engineering personnel, we may be unable to manage our growth or continue our technological development

Our continued growth in the provider business will depend upon our ability to attract and retain skilled employees, such as highly skilled nurses and other medical personnel. Competition for those employees is intense. Moreover, we believe that future success in the provider business will be significantly dependent on our ability to attract and retain qualified physicians to serve as medical directors of our dialysis clinics. Our dialysis products business depends on the development of new products, technologies and treatment concepts. Competition is also intense for skilled engineers and other technical research and development personnel. If we are unable to obtain the services of key personnel, the ability of our officers and key employees to manage our growth would suffer and our operations could suffer in other respects. These factors could preclude us from integrating acquired companies into our operations, which could increase our costs and prevent us from realizing synergies from acquisitions. Lack of skilled research and development personnel could impair our technological development, which would increase our costs and impair our reputation for production of technologically advanced products.

We face additional costs and uncertainties from international operations

We intend to expand our international presence. As a result, we expect that revenues from countries other than the U.S. and Germany will account for an increasing portion of future revenues.

Revenues from international operations are subject to a number of risks, including the following:

Fluctuations in exchange rates could adversely affect profitability;

We could face difficulties in enforcing and collecting accounts receivable under some countries' legal systems;

Local regulations could restrict our ability to obtain a direct ownership interest in dialysis clinics or other operations;

Political instability, especially in developing countries, could disrupt our operations;

Some customers and governments could have longer payment cycles, with resulting adverse effects on our cash flow; and

Some countries could impose additional taxes or restrict the import of our products.

The continuing financial crisis in Latin America and the decline of many of its major currencies against the U.S. dollar have affected our international operations and caused us to test our Latin America operations for impairment. See Item 5, Operating and Financial Review and Prospects-Financial Condition and Results of Operations. Any one or more of these factors, or any difficulty in integrating businesses we acquire into our operations, could increase our costs, reduce our revenues, or disrupt our operations, with possible material adverse effects on our business, financial condition and results of operations.

Other Risks

Our significant indebtedness may limit our ability to pay dividends or implement certain elements of our business strategy

We are substantially leveraged. As of December 31, 2003, our total consolidated liabilities were \$4.26 billion, including obligations with respect to all our trust preferred securities of approximately \$1.2 billion, our total consolidated assets were \$7.50 billion and our stockholders equity was \$3.24 billion. Our substantial level of debt presents the risk that we might not generate sufficient cash to service our indebtedness or that our

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leveraged capital structure could limit our ability to finance acquisitions and develop additional projects, to compete effectively or to operate successfully under adverse economic conditions.

Our senior credit agreement and the indentures relating to our trust preferred securities include covenants that require us to maintain certain financial ratios or meet other financial tests. Under our senior credit agreement, we are obligated to maintain a minimum consolidated net worth and a minimum consolidated fixed charge ratio (ratio of consolidated earnings before interest, taxes, depreciation, amortization and rent (EBITDAR) to fixed charges) and we are subject to a limit on our consolidated leverage ratio (ratio of consolidated funded debt to EBITDA).

Our senior credit agreement and our indentures include other covenants which, among other things, restrict or have the effect of restricting our ability to dispose of assets, incur debt, pay dividends, create liens or make capital expenditures, investments or acquisitions. These covenants may otherwise limit our activities. The breach of any of the covenants could result in a default under the credit agreement or the indentures, which could, in turn, create additional defaults under the agreements relating to our other long term indebtedness.

Because we are not organized under U.S. law, we are subject to certain less detailed disclosure requirements under U.S. federal securities laws

Under pooling agreements that we have entered into for the benefit of minority holders of our Ordinary shares and holders of our Preference shares (including, in each case, holders of American Depositary Receipts representing beneficial ownership of such shares), we have agreed to file quarterly reports with the Securities and Exchange Commission, to prepare annual and quarterly financial statements in accordance with U.S. generally accepted accounting principles, and to file information with the Securities and Exchange Commission with respect to annual and general meetings of our shareholders. However, we are a foreign private issuer, as defined in the Securities and Exchange Commission's regulations, and consequently we are not subject to all of the same disclosure requirements applicable to domestic companies. We are exempt from the Securities and Exchange Commission's proxy rules, and our annual reports contain less detailed disclosure than reports of domestic issuers regarding such matters as management, executive compensation and outstanding options, beneficial ownership of our securities and certain related party transactions. Also, our officers, directors and beneficial owners of more than 10% of our equity securities are exempt from the reporting requirements and short-swing profit recovery provisions of Section 16 of the Securities Exchange Act of 1934. We are also generally exempt from most of the governance rule revisions recently adopted by the New York Stock Exchange, other than the obligation to maintain an audit committee in accordance with Rule 10A-3 under the Securities Exchange Act of 1934, as amended. These limits on available information about our company may adversely affect the market prices for our securities.

Item 4. Information on the Company

A. History and Development of the Company

General

Fresenius Medical Care AG is a stock corporation (Aktiengesellschaft) organized under the laws of Germany. It was incorporated on August 5, 1996. Fresenius Medical Care AG is registered with the commercial register of the local court (*Amtsgericht*) of Hof an der Saale, Germany under HRB 2460. Our registered office (*Sitz*) is Hof an der Saale, Germany. Our business address is Else-Kröner-Strasse 1, 61352 Bad Homburg, Germany, telephone ++49-6172-609-0.

History

Fresenius Medical Care AG was created by the conversion of Sterilpharma GmbH, a limited liability company under German law organized in 1975, into a stock corporation under German law (*Aktiengesellschaft*). A shareholder's meeting on April 17, 1996 adopted the resolutions for this conversion and the commercial register registered the conversion on August 5, 1996.

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On September 30, 1996, we completed a series of transactions to consummate an Agreement and Plan of Reorganization entered into on February 4, 1996 by Fresenius AG and W.R. Grace which we refer to as the Merger elsewhere in this report. Pursuant to that agreement, Fresenius AG contributed Fresenius Worldwide Dialysis, its global dialysis business, including its controlling interest in Fresenius USA, Inc., in exchange for 35,210,000 Fresenius Medical Care Ordinary shares. Thereafter, we acquired:

all of the outstanding common stock of W.R. Grace, whose sole business at the time of the transaction consisted of National Medical Care, Inc., its global dialysis business, in exchange for 31,360,000 Ordinary shares; and

the publicly-held minority interest in Fresenius USA, in exchange for 3,430,000 Ordinary shares.

Effective October 1, 1996, we contributed all our shares in Fresenius USA to Fresenius Medical Care Holdings, which conducts business under the trade name Fresenius Medical Care North America, and which is the holding company for all of our operations in the U.S. and Canada and manufacturing operations in Mexico.

Capital Expenditures

We invested, by business segment and geographical areas, the following amounts during the three fiscal years ended December 31, 2003, 2002 and 2001 and have budgeted the following amounts for the year 2004:

	Actual (in millions)			Budget 2004
	2003	2002	2001	
Acquisitions				
North America	\$ 43	\$ 38	\$ 412	
International				
Germany	13			
Rest of World	45	50	49	
	<u> </u>	<u> </u>	<u> </u>	<u> </u>
Total Acquisitions	\$101	\$ 88	\$461	\$100
	<u> </u>	<u> </u>	<u> </u>	<u> </u>
Capital expenditures for property, plant and equipment				
North America	\$177	\$130	\$138	
International				
Germany	28	27	34	
Rest of World	86	82	103	
	<u> </u>	<u> </u>	<u> </u>	<u> </u>
Total Capital Expenditures	\$291	\$239	\$275	\$250
	<u> </u>	<u> </u>	<u> </u>	<u> </u>

During 2003, we finished the construction of three dialyzer assembly lines and two fiber spinning lines in our Ogden, Utah production facility. These expenditures were part of an effort to increase our single use dialyzer manufacturing capacity by 200% as part of our UltraCare program (see Business Overview Dialysis Care Fresenius UltraCare Program). In Asia Pacific we continued investment in production facilities. Other major 2003 capital expenditures were made for the improvement of production facilities in Germany, Italy and France. We finance our capital expenditures through cash flow from operations or under existing credit facilities.

In December 2003, we exercised an option to terminate an operating lease for certain manufacturing equipment in its Ogden, Utah, North American facility. The equipment was purchased for approximately \$66 million and is reflected as a capital expenditure in the accompanying consolidated statement of cash flows.

For information regarding recent acquisitions, see Business Overview Acquisitions.

B. Business Overview

Our Business

We are the world's largest kidney dialysis company engaged in both providing dialysis care and manufacturing dialysis products, based on publicly reported revenues and patients treated. We provide dialysis

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treatment to over 119,000 patients at our 1,560 clinics located in 25 countries. In the U.S. we also provide inpatient dialysis services, therapeutic apheresis, hemoperfusion and other services under contract to hospitals. We also develop and manufacture a complete range of equipment, systems and disposable products, which we sell to customers in over 100 countries. We use the information we gain when treating patients in developing new and improved products. We believe that our size, our activities in both dialysis care and dialysis products and our concentration in specific geographic areas allow us to operate more cost-effectively than many of our competitors. For the year ended December 31, 2003, we had net revenues of \$5.5 billion, an increase of 9% over 2002 revenues. We derived 70% of our revenues in 2003 from our North America operations and 30% from our International operations.

The following table summarizes net revenues for our North America segment and our International segment as well as our major categories of activity for the three years ended December 31, 2003, 2002 and 2001.

	<u>2003</u>	<u>2002</u>	<u>2001</u>
	(in millions)		
North America			
Dialysis Care	\$ 3,429	\$ 3,294	\$ 3,131
Dialysis Products ⁽¹⁾	426	454	471
	<u>3,855</u>	<u>3,748</u>	<u>3,602</u>
International			
Dialysis Care	550	415	426
Dialysis Products	1,123	921	831
	<u>1,673</u>	<u>1,336</u>	<u>1,257</u>

(1) We evaluate North America product sales based on net available external market. See Item 5.A. Operating Results for explanation and analysis.

Renal Industry Overview**End-Stage Renal Disease**

End-stage renal disease (ESRD) is the stage of advanced chronic kidney disease that is characterized by the irreversible loss of kidney function and requires regular dialysis treatment or kidney transplantation to sustain life. A normally functioning human kidney removes waste products and excess water from the blood, which prevents toxin buildup, water overload and the eventual poisoning of the body. A number of conditions — diabetes, hypertension, glomerulonephritis and inherited diseases — can cause chronic kidney disease. Nearly 60% of all people with ESRD acquire the disease as a complication of one or more of these primary conditions.

There are currently only two methods for treating ESRD: dialysis and kidney transplantation. Scarcity of compatible kidneys limits transplants. According to data published by the Centers for Medicare and Medicaid Services (CMS) (formerly the Health Care Financing Administration) of the U.S. Department of Health and Human Services, 14,628 patients of the ESRD patient population, received kidney transplants in the U.S. during 2001 an increase of 2% over 2000. According to the United States Renal Data System (USRDS) 2002 Annual Report only 2 - 3% of all incident patients received a pre-emptive transplant in 2000. In Germany, the third biggest dialysis market worldwide according to our own internal survey, less than 1% of all incident patients received pre-emptive transplants, as published by the German registry Quasi-Niere. Therefore, most patients suffering from ESRD must rely on dialysis, which is the removal of toxic waste products and excess fluids from the body by artificial means. There are two major dialysis methods commonly used today, hemodialysis (HD) and peritoneal dialysis (PD). These are described below under Treatment Options for ESRD. Generally, an ESRD patient's physician, in consultation with the patient, chooses the patient treatment method, which is based on the patient's medical conditions and needs.

Based on data published by the CMS, the number of patients in the U.S. who received dialysis for chronic ESRD grew from approximately 66,000 in 1982 to 285,982 at December 31, 2001, a compound annual rate of 8%. We believe that growth will continue at 4 - 6% per year. According to data from our own internal survey, the

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number of non-U.S. chronic dialysis patients is growing at estimated annual rates of 7 - 8% for patients receiving dialysis. At the end of 2002, an estimated 1.2 million patients were undergoing dialysis treatment worldwide. According to our own market surveys, Japan is the second largest dialysis market in the world. According to data published by the Japanese Society for Dialysis Therapy, more than 200,000 dialysis patients were being treated at the end of 2000. In the rest of the world, we estimate that at the end of 2002 there were approximately 292,000 dialysis patients in Europe, approximately 160,000 patients in Asia (excluding Japan) and approximately 143,000 patients in Latin America. We believe that the continuing growth in the number of dialysis patients is principally attributable to:

increased general life expectancy and the overall aging of the general U.S. and European population;

shortage of donor organs for kidney transplants;

improved dialysis technology that has expanded the patient population able to undergo life-prolonging dialysis;

greater access to treatment in developing countries.

better treatment and survival of patients with hypertension, diabetes and other illnesses that lead to ESRD.

Treatment Options for ESRD

Hemodialysis. Hemodialysis removes toxins and excess fluids from the blood whereby the blood flows outside the body through plastic tubes known as bloodlines into a specially designed filter, called a dialyzer. The dialyzer functions as an artificial kidney by separating waste products and excess water from the blood. Dialysis solution flowing through the dialyzer carries away the waste products and excess water, and supplements the blood with solutes that have been depleted due to renal failure. The treated blood is returned to the patient. The hemodialysis machine pumps blood, adds anti-coagulants, regulates the purification process and controls the mixing of dialysis solution and the rate of its flow through the system. This machine can also monitor and record the patient's vital signs.

Hemodialysis patients generally receive treatment three times per week, typically for around three to five hours per treatment. The majority of hemodialysis patients receive treatment at outpatient dialysis clinics, such as ours, where hemodialysis treatments are performed with the assistance of a nurse or dialysis technician under the general supervision of a physician.

According to data from the CMS, as of December 31, 2001, there were 4,081 Medicare-certified ESRD treatment clinics in the U.S. Ownership of these clinics is fragmented. We estimate that there were approximately 4,900 dialysis clinics in Europe at the end of 2002, of which almost 60% are government-owned, more than 30% are privately owned, and around 10% are operated by health care organizations. In Latin America, privately owned clinics predominate, comprising over 70% of all clinics providing dialysis care.

According to the CMS, as of December 31, 2001, hemodialysis patients represented about 90% of all dialysis patients in the U.S. Our most recent studies indicate hemodialysis patients comprise 95% of all dialysis patients in Japan, 89% in the European Union and 85% in the rest of the world.

Peritoneal Dialysis. Peritoneal dialysis removes toxins from the blood using the peritoneum, the membrane lining covering the internal organs located in the abdominal area. Peritoneal dialysis patients administer their own treatments in their own homes and workplaces, either by a treatment known as continuous ambulatory peritoneal dialysis or CAPD, or by a treatment we introduced in 1980 known as continuous cycling peritoneal dialysis or CCPD. In both of these treatments, a surgically implanted catheter provides access to the peritoneal cavity. Using this catheter, the patient introduces a sterile dialysis solution from a solution bag through a tube into the peritoneal cavity. The peritoneum operates as the filtering membrane and, after a specified dwell time, the solution is drained and disposed of. A typical CAPD peritoneal dialysis program involves the introduction and disposal of dialysis solution four times a day. With CCPD a machine cycles solution to and from the patient's peritoneal cavity while the patient sleeps.

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Our Strategy

Our objective is generating revenue growth that exceeds market growth of the dialysis industry, measured by growth in the patient population, while maintaining our leading position in the market and increasing earnings at a faster pace than revenues. Our dialysis care and product revenues have grown faster than the market over the past five years, and we believe that we are well positioned to meet our objectives by focusing on the following strategies:

Continue to Provide High Standards of Patient Care. We believe that our reputation for providing the highest standards of patient care is a competitive advantage.

Differentiated Patient Care Programs from those of Our Competitors. We believe that our UltraCare Patient Care program offered at our North America dialysis facilities will distinguish and differentiate our patient care programs from those of our competitors. UltraCare therapy employs single-use high flux polysulfone dialyzers, on-line quality measurement, and Ultra Pure Dialysate, all of which we feel improves mortality and increases the quality of patient care. The change to single-use dialyzers has increased our per treatment dialyzer costs relative to use of multi-use dialyzers. These cost increases have been offset, however, by our ability to achieve economies of scale in the production of these dialyzers, due to our large-scale single-use dialyzer manufacturing capacity. Moreover, we have implemented a new staffing model based on single-use that reduces our personnel costs per treatment. Finally, automated controls in our new 2008 Series dialysis machine reduces concentrate usage and associated costs.

Expand Presence in Attractive Growth Markets Worldwide. We intend to continue to take advantage of the reputation and market recognition our global product business has created by acquiring and establishing new dialysis clinics within attractive international markets. We believe that we will obtain an increasing percentage of our dialysis care growth from worldwide markets. We believe that increases in per capita income in developing countries will make general health care benefits, which may include payment for dialysis treatment, more widely available and present significant opportunities.

Increase Our Spectrum of Dialysis Services. One of our objectives is to continue to expand our role within the broad spectrum of services for dialysis patients. We have begun to implement this strategy by providing expanded and enhanced patient services, including laboratory services, to both our own clinics and those of third parties. We estimate that our Spectra Renal Management division provides laboratory services for approximately 39% of the ESRD patients in the U.S. We have developed disease state management methodologies, which involve the coordination of total patient care for ESRD patients and which we believe are attractive to managed care payors. We have formed Optimal Renal Care, LLC, a joint venture with The Permanente Federation. We also formed Renaissance Health Care as a joint venture with participating nephrologists. Renaissance provides ESRD and Chronic Kidney Disease programs to more than 3,000 patients. We also operate a surgical center for the management and care of vascular access for patients which decreases hospitalization.

Offer Complete Dialysis Product Lines with Recurring Disposable Products Revenue Streams. We offer broad and competitive hemodialysis and peritoneal dialysis product lines. These product lines enjoy broad market acceptance and enable us to serve as our customers' single source for all of their dialysis machines, systems and disposable products.

Extend Our Position as an Innovator in Product and Process Technology. We are committed to technological leadership in both hemodialysis and peritoneal dialysis products. We have an over 388 member research and development team that focuses on developing dialysis systems that are safer, more effective and easier to use and that can be easily customized to meet the differing needs of customers around the world. We believe that our extensive expertise in patient treatment and clinical data will further enhance our ability to develop more effective products and treatment methodologies. Our ability to manufacture dialysis products on a cost-effective and competitive basis results in large part from our process technologies. Over the past several years, we have reduced manufacturing costs per unit through development of proprietary manufacturing technologies that have streamlined and automated our production processes.

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Dialysis Care

Dialysis Services

We provide dialysis treatment and related laboratory and diagnostic services at our approximately 1,560 outpatient dialysis clinics, 1,110 of which are in the U.S. and 450 of which are in 23 countries outside of the U.S. Our operations outside the U.S. generated 14% of our 2003 dialysis care revenue. We currently operate dialysis clinics in Argentina, Australia, Brazil, China, Colombia, Chile, Czech Republic, France, Germany, Hungary, Hong Kong, Italy, Singapore, Mexico, Portugal, Poland, Slovakia, Slovenia, South Africa, Spain, Taiwan, Turkey, United Kingdom and Venezuela. Our dialysis clinics are generally concentrated in areas of high population density. In 2003, we acquired 42 existing clinics, opened 76 new clinics and consolidated 38 clinics. The number of patients we treat at our clinics increased by about 6%, from approximately 112,200 at December 31, 2002 to approximately 119,250 at December 31, 2003.

With our large patient population, we have developed proprietary patient statistical databases which enable us to improve dialysis treatment outcomes, and improve the quality and effectiveness of dialysis products. We believe that local physicians, hospitals and managed care plans refer their ESRD patients to our clinics for treatment due to:

our reputation for quality patient care and treatment;

our extensive network of dialysis clinics, which enables physicians to refer their patients to conveniently located clinics; and

our reputation for technologically advanced products for dialysis treatment.

We treat approximately 26% of the dialysis patients in the U.S. including those patients treated in clinics we manage. Based on publicly available reports, we believe our next largest competitor treats approximately 15% of U.S. dialysis patients. For the year 2003, dialysis services accounted for 72% of our total revenue.

At our clinics, we provide hemodialysis treatments at individual stations through the use of dialysis machines and disposable products. A nurse attaches the necessary tubing to the patient and the dialysis machine and monitors the dialysis equipment and the patient's vital signs. The capacity of a clinic is a function of the number of stations and such factors as type of treatment, patient requirements, length of time per treatment, and local operating practices and ordinances regulating hours of operation.

Each of our dialysis clinics is under the general supervision of a Medical Director and, in some cases, one or more associate Medical Directors, all of whom are physicians. See Patients, Physician and Other Relationships. Each dialysis clinic also has an administrator or clinical manager who supervises the day-to-day operations of the facility and the staff. The staff typically consists of registered nurses, licensed practical nurses, patient care technicians, a social worker, a registered dietician, a unit clerk and biomedical technicians.

As part of the dialysis therapy, we provide a variety of services to ESRD patients in the U.S. at our dialysis clinics. These services include administering EPO, a bioengineered protein that stimulates the production of red blood cells. EPO is used to treat anemia, a medical complication that ESRD patients frequently experience, and we administer EPO to most of our patients. Revenues from EPO accounted for approximately 23% of dialysis care revenue in our North America segment for the year ended December 31, 2003. We receive a substantial majority of this revenue as reimbursements through the Medicare and Medicaid programs. Amgen Inc., is the sole manufacturer of EPO in North America, and any interruption of supply could materially adversely affect our business, financial condition and results of operations. Our current contract with Amgen covers the period from January 2004 to January 2006.

Our clinics also offer services for home dialysis patients, the majority of whom receive peritoneal dialysis treatment. For those patients, we provide materials, training and patient support services, including clinical monitoring, follow-up assistance and arranging for delivery of the supplies to the patient's residence. In the U.S. clinic services include the supplying of EPO. See Regulatory and Legal Matters Reimbursement U.S. for a discussion of billing for these products and services.

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We also provide dialysis services under contract to hospitals in the U.S. on an "as needed" basis for hospitalized ESRD patients and for patients suffering from acute kidney failure. Acute kidney failure can result from trauma or similar causes, and requires dialysis until the patient's kidneys recover their normal function. We service these patients either at their bedside, using portable dialysis equipment, or at the hospital's dialysis site. Contracts with hospitals provide for payment at negotiated rates that are generally higher than the Medicare reimbursement rates for chronic in-clinic out-patient treatments.

We employ a centralized approach with respect to certain administrative functions common to our operations. For example, each dialysis clinic uses our proprietary manuals containing our standardized operating and billing procedures. We believe that centralizing and standardizing these functions enhance our ability to perform services on a cost-effective basis.

The manner in which each clinic conducts its business depends, in large part, upon applicable laws, rules and regulations of the jurisdiction in which the clinic is located, as well as our clinical policies. However, a patient's attending physician, who may be the clinic's Medical Director or an unaffiliated physician with staff privileges at the clinic, has medical discretion to prescribe the particular treatment modality and medications for that patient. Similarly, the attending physician has discretion in prescribing particular medical products, although the clinic typically purchases equipment, regardless of brand, in consultation with the Medical Director.

Fresenius UltraCare Program

In 2002, we started a new program in the North America dialysis services group called UltraCare. This program combines our latest product technology with our highly trained and skilled staff to offer our patients a superior level of care. The basis for this form of treatment is the Optiflux polysulfone single-use dialyzer. These dialyzers have excellent blood detoxification properties and are the most efficient dialyzers currently available. Optiflux dialyzers are combined with our 2008 Hemodialysis Delivery System series, which has advanced online patient monitoring as well as several systems to allow the tailoring of treatment to meet individual patient needs. Among the other capabilities of this system, staff will be alerted if toxin clearance is less than the target prescribed for the patient, and treatment can be adjusted accordingly.

Laboratory Services

We provide laboratory testing and marketing services through Spectra Renal Management. Spectra Renal Management is the leading U.S. dialysis clinical laboratory providing blood, urine and other bodily fluid testing services to assist physicians in determining whether a dialysis patient's therapy regimen, diet and medicines remain optimal. Spectra Renal Management operates two laboratories, located in New Jersey and Northern California. During the year ended December 31, 2003, Spectra Renal Management performed over 38 million tests for more than 120,000 dialysis patients in 1,758 clinics across the U.S. including clinics that we do own or operate.

Acquisitions

A significant factor in the growth in our revenue and operating earnings in prior years has been our ability to acquire health care businesses, particularly dialysis clinics, on reasonable terms. Worldwide, physicians own many dialysis clinics that are potential acquisition candidates for us. In the U.S., doctors might determine to sell their clinics to obtain relief from day-to-day administrative responsibilities and changing governmental regulations, to focus on patient care and to realize a return on their investment. Outside of the U.S., doctors might determine to sell and/or enter into joint ventures or other relationships to us to achieve the same goals and to gain a partner with extensive expertise in dialysis products and services.

We paid aggregate cash consideration for acquisitions of Fresenius AG's adsorber business and new clinics of approximately \$92 million in 2003 and approximately \$80 million in 2002, primarily for dialysis clinics. In 2003, we completed new acquisitions and acquisitions of previously managed clinics totaling 42 dialysis facilities. These acquisitions expand our presence in selected key geographic areas.

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We continued to enhance our presence in the U.S. and abroad by acquiring individual or small groups of dialysis clinics in selected markets, expanding existing clinics, and opening new clinics.

Quality Assurance in Dialysis Care

Beginning in 2001, our quality management activities were primarily focused on comprehensive development and implementation of an Integrated Quality Management System (IMS). Our goals in this area included not only meeting quality requirements for our dialysis clinics and environmental concerns, but also managing the quality of our dialysis care. This approach resulted in an IMS structure that closely reflects existing corporate processes. We also were able to use the IMS to fulfill many legal and normative regulations covering service lines. In addition, the IMS offers a highly flexible structure that allows us to adapt to future regulations. The most important of these include, among others, ISO 9001, EN 46001, ISO 13485, and 21 CFR 820 which establish quality control and other performance criteria.

In 2003, internal and external auditors inspected our dialysis clinics. They confirmed the effectiveness of our organization and processes and documented our compliance with relevant regulations. The use of newly developed evaluation methods allowing simpler performance comparisons identified additional improvement possibilities. Another focus of our activities in 2003 was the continuing certification of our dialysis clinics under ISO 9001, EN 46001, ISO 13485, and 21 CFR 820 .

The rapid pace of IMS integration will continue in 2004. The integration of a new risk and complaint management system and the further involvement of our subsidiaries in the Asian-Pacific and Latin American regions are additional goals.

At each of our North America dialysis clinics, a quality assurance committee is responsible for reviewing quality of care data, setting goals for quality enhancement and monitoring the progress of quality assurance initiatives. We believe that we enjoy a reputation of providing high quality care to dialysis patients. In 2003, the Company continued to develop and implement programs to assist in achieving our quality goals. Our Access Intervention Management Program (AIM), started in 2001, detects and corrects arteriovenous access failure in hemodialysis treatment, which is the major cause of hospitalization and morbidity.

Sources of U.S. Dialysis Care Net Revenue

The following table provides information for the years ended December 31, 2003, 2002 and 2001 regarding the percentage of our U.S. dialysis treatment services net revenues from (a) the Medicare ESRD program, (b) private/ alternative payors, such as commercial insurance and private funds, (c) Medicaid and other government sources and (d) hospitals.

	Year Ended December 31,		
	2003	2002	2001
Medicare ESRD program	61.0%	61.5%	59.7%
Private/alternative payors	29.2%	29.5%	31.2%
Medicaid and other government sources	3.9%	4.5%	4.6%
Hospitals	5.9%	4.5%	4.5%
Total	100.0%	100.0%	100.0%

Under the Medicare ESRD program, Medicare reimburses dialysis providers for the treatment of certain individuals who are diagnosed as having ESRD, regardless of age or financial circumstances. See Regulatory and Legal Matters Reimbursement.

Patient, Physician and Other Relationships

We believe that our success in establishing and maintaining dialysis clinics, both in the U.S. and in other countries, depends significantly on our ability to obtain the acceptance of and referrals from local physicians,

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hospitals and managed care plans. A dialysis patient generally seeks treatment at a conveniently located clinic at which the patient's nephrologist has staff privileges.

Medicare ESRD program reimbursement regulations require that a Medical Director generally supervise treatment at a dialysis clinic. Generally, the Medical Director must be board certified or board eligible in internal medicine and have at least twelve months of training or experience in the care of patients at ESRD clinics. Our Medical Directors also maintain their own private practices.

Competition

Dialysis Services. The dialysis services industry is highly competitive. Our major competitors in dialysis services include Gambro AB, DaVita, Inc. (formerly Total Renal Care), Baxter International Inc., Renal Care Group and the Kuratorium für Dialyse und Nierentransplantation e.V. Ownership of dialysis clinics in the U.S. is fragmented with a large number of operators each owning 10 or fewer clinics and a small number of larger multi-clinic providers, of which we are the largest. Industry consolidation has been ongoing over the last decade. Many of our dialysis clinics are in urban areas, where there frequently are many competing clinics in proximity to our clinics. We experience direct competition from time to time from former Medical Directors, former employees or referring physicians who establish their own clinics. Furthermore, other health care providers or product manufacturers, some of who have significant operations, may decide to enter the dialysis business in the future.

Because in the U.S. government programs are the primary source of reimbursement for services to the majority of patients, competition for patients in the U.S. is based primarily on quality and accessibility of service and the ability to obtain admissions from physicians with privileges at the facilities. However, the extension of periods during which commercial insurers are primarily responsible for reimbursement and the growth of managed care have placed greater emphasis on service costs for patients insured with private insurance.

In most countries other than the U.S., we compete primarily against individual free-standing clinics and hospital-based clinics. In many of these countries, especially the developed countries, governments directly or indirectly regulate prices and the opening of new clinics. Providers compete in all countries primarily on the basis of quality and availability of service and the development and maintenance of relationships with referring physicians.

Laboratory Services. Spectra Renal Management competes in the U.S. with large nationwide laboratories, dedicated dialysis laboratories and numerous local and regional laboratories, including hospital laboratories. In the laboratory services market, companies compete on the basis of performance, including quality of laboratory testing, timeliness of reporting test results and cost-effectiveness. We believe that our services are competitive in these areas.

Dialysis Products

We are currently the world's largest manufacturer and distributor of equipment and related products for hemodialysis and the second largest manufacturer of peritoneal dialysis products, based on publicly reported revenues, with operations in Germany, the U.S., and in 35 other countries. We sell our dialysis products directly and through distributors in over 100 countries. Most of our customers are dialysis clinics. For the year 2003, dialysis products accounted for 28% of our total revenue.

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The following table shows the breakdown of our dialysis product revenues into sales of hemodialysis products and peritoneal dialysis products.

	Year Ended December 31,					
	2003		2002		2001	
	Total Product Revenues	% of Total	Total Product Revenues	% of Total	Total Product Revenues	% of Total
	(U.S. dollars in millions)					
Hemodialysis Products	\$ 1,326.1	86	\$ 1,181.0	86	\$ 1,114.0	86
Peritoneal Dialysis Products	211.5	14	194.2	14	188.2	14
Total	\$ 1,537.6	100	\$ 1,375.2	100	\$ 1,302.2	100

Hemodialysis Products

We offer a comprehensive hemodialysis product line and believe that our broad range of technologically sophisticated hemodialysis products makes us a leader in the hemodialysis product field. We continually strive to expand and improve the capabilities of our hemodialysis systems to offer an advanced treatment mode at reasonable cost.

Dialysis Machines. We sell our dialysis machines as Series 2008H and 2008K models in North America and Series 4008 models in the rest of the world. Our dialysis machines offer the following features and advantages:

Volumetric dialysate balancing and ultrafiltration control system. This system, which we introduced in 1977, provides for safe and more efficient use of highly permeable dialyzers, permitting faster dialysis with controlled rates of fluid removal;

Proven hydraulic systems, providing reliable operation and servicing flexibility;

Compatibility with all manufacturers' dialyzers and a wide variety of blood-lines and dialysis solutions, permitting maximum flexibility in both treatment and disposable products usage;

Modular design, which permits us to offer dialysis clinics a broad range of options to meet specific patient or regional treatment requirements. Modular design also allows upgrading through module substitution without replacing the entire machine;

Specialized modules that provide monitoring and response capability for selected bio-physical patient parameters, such as body temperature, relative blood volume and electrolyte balances. This concept, known as physiological dialysis, permits hemodialysis treatments with lower incidence of a variety of symptoms or side effects, which still occur frequently in standard hemodialysis.

Sophisticated microprocessor controls, and display and readout panels that are adaptable to meet local language requirements;

Battery backup, which continues operation of the blood circuit and all protective systems up to 20 minutes following a power failure;

Online clearance, measurement of dialyzer clearance for quality assurance with the On-Line Clearance Monitor, providing immediate effective clearance information, real time treatment outcome monitoring, and therapy adjustment during dialysis without requiring invasive procedures or blood samples;

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On-line data collection capabilities and computer interfacing with our FINESSE module and FDS08® system. Our machines can:

monitor and assess prescribed therapy;

connect a large number of hemodialysis machines and peripheral devices, such as patient scales, blood chemistry analyzers and blood pressure monitors, to a personal computer network;

enter nursing records automatically at bedside to register and document patient treatment records, facilitate billing, and improve record-keeping and staff efficiency;

adapt to new data processing devices and trends;

perform home hemodialysis with remote monitoring by a staff caregiver; and

record and analyze trends in medical outcome factors in hemodialysis patients.

Dialyzers. We manufacture dialyzers using hollow fiber polysulfone membranes, a synthetic material. We are the leading worldwide producer of polysulfone dialyzers. We believe that polysulfone offers the following superior performance characteristics compared to other materials used in dialyzers:

higher biological compatibility, resulting in reduced incidence of adverse reactions to the fibers;

greater capacity to clear uremic toxins from patient blood during dialysis, permitting more thorough, more rapid dialysis, resulting in shorter treatment time; and

a complete range of permeability, or membrane pore size, which permits dialysis at prescribed rates – high flux, medium flux and low flux, as well as ultra flux for acute dialysis, and allows tailoring of dialysis therapy to individual patients.

Single Use Dialyzers. In North America, we have completed a \$65 million capital investment program to significantly expand the capacity of our dialyzer manufacturing plant in Ogden, Utah through the addition of three new dialyzer assembly lines.

Other Hemodialysis Products.

We manufacture and distribute arterial, venous, single needle and pediatric bloodlines. We produce both liquid and dry dialysate concentrates. Liquid dialysate concentrate is mixed with purified water by the hemodialysis machine to produce dialysis solution, which removes the toxins and excess water from the patient's blood during dialysis. Dry concentrate, developed more recently, is less labor-intensive to use, requires less storage space and may be less prone to bacterial growth than liquid solutions. We also produce dialysis solutions in bags, including solutions for priming and rinsing hemodialysis bloodlines, as well as connection systems for central concentrate supplies and devices for mixing dialysis solutions and supplying them to hemodialysis machines. Other products include solutions for disinfecting and decalcifying hemodialysis machines, fistula needles, hemodialysis catheters, and products for acute renal treatment.

Peritoneal Dialysis Products

We offer a full line of peritoneal dialysis products. We manufacture peritoneal dialysis solutions in bags, peritoneal dialysis cycling machines for CCPD and disposable products for both CAPD and CCPD, such as tubing, sterile solutions and sterile kits to prepare patients for dialysis. We also distribute other manufacturers' peritoneal dialysis products, primarily to our own dialysis clinics.

CAPD Systems. We manufacture standard and specialized peritoneal dialysis solutions. We believe that our peritoneal dialysis products offer significant advantages for CAPD, including:

ease of use and greater protection against contamination by touch than other peritoneal dialysis systems presently available. Our products incorporate our Safe-Lock connection system for introducing and draining dialysis solution into and from the abdominal cavity. Our A.N.D.Y. and A.N.D.Y. Plus systems,

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which include a special drainage bag and a snap-off Y-shaped piece connected to the Safe-Lock connector at the catheter, provide protection from contamination in a dual-bag system;

suitability for all peritoneal dialysis patients through the Inpersol and Safe-Lock product lines. Inpersol products are interchangeable with those of other manufacturers; Safe-Lock products may be used only by peritoneal dialysis patients whose catheters include the Safe-Lock connector, which attaches to a solution bag fitted with the other part;

manufacture with Biofine, a new environmentally friendly, more biocompatible, plastic material for foils, tubings and other parts of peritoneal dialysis systems.

the benefits of Biofine with protection against contamination through our new Stay-Safe system, launched in 1997. The system comprises tubing, connectors and a peritoneal dialysis solution double bag, made entirely from Biofine. It uses a single switching mechanism that replaces three tubing clamps to control solution drainage, flushing of tubes that connect solution bags to catheters, and introduction of new solution. The single switch also provides tight closure of the line and, to further reduce the possibility of contamination, the switch seals catheter access and surrounds the catheter adapter with disinfectant;

higher solution bag volumes with our Premier twin-bag system which provides solution container and pre-attached tubing set in one package. The higher solution volumes permit larger dosages without increasing the number of required daily solution exchanges performed by the patient; and

improved biocompatibility with CAPD stay.safe Balance, a lactate-buffered peritoneal dialysis solution that has a pH balance in the human physiological range.

CCPD Products. We introduced the first peritoneal dialysis cyclor machine in 1980. In a standard CAPD program, a patient manually introduces two liters of fresh peritoneal dialysis solution and drains the used solution four times over a 24-hour period. Treatment occurs seven days per week and the patient must perform the treatment while awake. With CCPD therapy, the cyclor automatically delivers a prescribed volume of dialysis solution into the peritoneal cavity through an implanted catheter, allows the solution to dwell for a specified time, and completes the process by draining the solution. CCPD therapy offers the following benefits over CAPD:

Solution exchanges take place automatically, which may reduce the risk of peritonitis due to less frequent handling of the catheter and connections;

The patient can cycle at home, throughout the night while asleep. The patient has complete daytime freedom, wearing only the surgically implanted catheter and capping device; and

CCPD delivers more effective therapy than CAPD due to the supine position of the patient during the night, higher volume exchanges and preferable cycle management which can be particularly significant for patients who need more therapy due to body size, ultrafiltration loss or other reasons.

Our cycling equipment incorporates microprocessor technology, and the patient, hospital or clinic staff can easily program it to perform specific prescribed therapy for a given patient. Since all components are monitored and programmable, these machines allow the physician to prescribe any of a number of current therapy procedures. Our CCPD products and therapies include:

the Sleep-Safe Cyclor, a new cyclor with an extremely compact and light design, that we began marketing in late 1999. Its pumping mechanism and disposable cartridge allow exact delivery of the peritoneal dialysis solution;

PD-PLUS, a variant on CCPD therapy we introduced in 1994. PD-PLUS therapy provides a more tailored therapy than regular CCPD using a simpler nighttime cyclor and, where necessary, includes one manual dialysis solution exchange during the day. We believe that PD-PLUS therapy is less costly and easier to administer than typical CCPD. We also believe that PD-PLUS therapy improves toxin removal by more than 40% compared to CAPD. By increasing the effectiveness of peritoneal dialysis treatments, PD-PLUS may also effectively prolong the time period during which a patient will be able to remain on

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peritoneal dialysis before requiring hemodialysis. PD-PLUS therapy can only be performed using the Fresenius Freedom Cyclers and special tubing using Safe-Lock connectors; and

IQcard, for use with the Freedom Cyclers PD-PLUS to monitor CCPD therapy for a full treatment history and improved therapy compliance.

Other Peritoneal Dialysis Products. We also manufacture and distribute pediatric treatment systems for administration of low volumes of dialysis solutions, assist devices to facilitate automated bag exchange for handicapped patients, catheters, catheter implantation instruments, silicon glue, Pack-PD, a computer program which analyzes patient and peritoneal characteristics to present a range of treatment options for individual therapies, disinfectants, bag heating plates adapters, and products to assist and enhance connector sterility. We also provide scientific and patient information products, including support materials, such as brochures, slides, videos, instructional posters and training manuals.

Marketing, Distribution and Service

We sell most of our products to clinics, hospitals and specialized treatment clinics. With our comprehensive product line and years of experience in dialysis, we believe that we have been able to establish and maintain very close relationships with our clinic customer base on a global basis. Close interaction between our sales force and research and development personnel enables us to integrate concepts and ideas that originate in the field into product development. We maintain a direct sales force of trained salespersons engaged in the sale of both hemodialysis and peritoneal dialysis products. This sales force engages in direct promotional efforts, including visits to physicians, clinical specialists, hospitals, clinics and dialysis clinics, and represents us at industry trade shows. We also sponsor medical conferences and scientific symposia as a means for disseminating product information. Our clinical nurses provide clinical support, training and assistance to customers and assist our sales force. We also use outside distributors to provide sales coverage in countries that our internal sales force does not service.

In our basic distribution system, we ship products from factories to central warehouses which are frequently located near the factories. From this central warehouse, we distribute our dialysis products to regional warehouses. We then distribute peritoneal dialysis products to the patient at home, and ship hemodialysis products directly to dialysis clinics and other customers. Local sales forces, independent distributors, dealers and sales agents sell all our products.

We offer customer service, training and education in the applicable local language, and technical support such as field service, repair shops, maintenance, and warranty regulation for each country in which we sell dialysis products. We provide training sessions on our equipment at our facilities in Schweinfurt, Germany, Chicago, Illinois and Walnut Creek, California and we also maintain regional service centers that are responsible for day-to-day international service support.

Manufacturing Operations

We operate state-of-the-art production facilities world wide to meet the demand for machines, cyclers, dialyzers, solutions, concentrates, mixes, bloodlines, and disposable tubing assemblies and equipment for water treatment in dialysis clinics. We have invested significantly in developing proprietary processes, technologies and manufacturing equipment which we believe provide a competitive advantage in manufacturing our products. We are using our facilities in St. Wendel, Germany and Ogden, Utah as centers of competence for development and manufacturing and to implement similar technologies at our other facilities.

We produce and assemble hemodialysis machines and CCPD cyclers in our Schweinfurt, Germany and our Walnut Creek, California facilities. We also maintain facilities at our service and local distribution centers in Argentina, Egypt, France, Italy, The Netherlands, China, Brazil and Russia for testing and calibrating dialysis machines manufactured or assembled elsewhere, to meet local end user market needs. We manufacture and assemble dialyzers and polysulfone membranes in our St. Wendel, Germany, L Arbresle, France and Inukai, Japan facilities and at production facilities of our joint ventures in Belarus, Saudi Arabia and Japan. At our Ogden, Utah facilities we manufacture and assemble dialyzers and polysulfone membranes as well as

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manufacture PD solutions. We have opened PD production in Mexico. Our facilities are inspected on a regular basis by national and/or international authorities.

In North America we expanded our manufacturing capacity substantially. During 2003, we completed a \$65 million capital commitment to significantly expand the capacity of our dialyzer manufacturing plant in Ogden, Utah through the addition of three new dialyzer assembly lines. See *History and Development of the Company* *Capital Expenditures*.

Sources of Supply

Our purchasing policy combines worldwide sourcing of high-quality materials with the establishment of long-term relationships with our suppliers. Additionally, we carefully assess the reliability of all materials purchased to ensure that they comply with the rigorous quality and safety standards required for our dialysis products. Our International Purchasing Consulting Center (PCC) ensures that we consistently maintain high standards by entering into global agreements. An interactive information system links all our global projects to ensure that they are standardized and constantly monitored.

PCC focuses on further optimizing procurement logistics and reducing purchasing costs. Supplemental raw material contracts for all manufacturers of semi-finished goods will enable us to improve purchasing terms for our complete network. We will also intensify our use of internet-based procurement tools by purchasing raw materials through special on-line auctions. Our sophisticated routing software enables us to distribute our supplies to best accommodate customer requests while maintaining operational efficiency.

New Product Introductions

Research and development focuses strongly on the development of new products, technologies and treatment concepts to optimize treatment quality for dialysis patients, and on process technology for manufacturing our products. Research and development expenditures were \$50 million in 2003, \$47 million in 2002, and \$36 million in 2001.

New or enhanced products introduced in 2003 included the following:

Patient Online (POL). A software tool to manage PD therapy.

MultiBic. A dialysis substitution fluid registered and introduced this year in Germany.

Patents, Trademarks and Licenses

As the owner of or licensee under patents and trademarks throughout the world, we hold rights under about 1,100 patents and patent applications relating to dialysis technology in major markets. Patented technologies that relate to dialyzers include our polysulfone hollow fiber, an in-line sterilization method, and sterile closures for in-line sterilized medical devices. The more recent generation of DIASAFEplus filters and FX dialyzers are also the subject of patents and pending patent applications.

The Company holds the exclusive license on European patents/patent applications on the Autoprime technology for the automated priming of the extracorporeal hemodialysis blood circuit with dialyzing liquid through the membrane of the dialyzer.

The connector system for our biBag bicarbonate concentrate powder container has been patented in the USA, Norway and Europe while national applications in Japan and Finland are still pending.

Among the Company's more significant protective rights, one patent family protects the Company's polysulfone hollow fiber until 2007 in the United States, and until 2005 in other main markets. The in-line sterilization method is patented until 2010 and the biBag connector is protected until 2013, both in Germany, in the United States, and in other important markets. The dates given represent the maximum life time of the corresponding patents. The Company believes that even after expiration of these patents, our proprietary know-how for the manufacture of these products will continue to constitute a competitive advantage.

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For peritoneal dialysis, the Company holds protective rights on our polyolefine film Biofine, suitable for packaging intravenous and peritoneal dialysis fluids. These patents have been granted in Australia, Germany, and the USA, with patent applications pending in various other countries. A further pending patent family describes a special film for a peelable, non-PVC, multi chamber bag for peritoneal dialysis solutions. A U.S. patent has already been granted.

We believe that our success will depend, in large part, on our technology. As a standard practice, we obtain legal protections we believe are appropriate for our intellectual property. Intellectual property is, however, subject to infringement or invalidation claims. In addition, technological developments in ESRD therapy could reduce the value of our existing intellectual property. Any such reduction could be rapid and unanticipated. Other than as disclosed in this report, we are not dependent to any material extent upon patents, licenses or contracts.

Competition

The markets in which we sell our dialysis products are highly competitive. Our competitors in the sale of hemodialysis and peritoneal dialysis products include Gambro AB, Baxter International, Inc., Asahi Medical Co., Ltd., Bellco S.p.A., a subsidiary of Sorin Biomedica S.p.A., Bieffe Medital S.p.A., which is an affiliate of Baxter International, Inc., B. Braun Melsungen AG, Nisscho Corporation, including Nisscho Nipro Corporation Ltd., Nikkiso Co., Ltd., Terumo Medical Corporation and Toray Medical Co., Ltd. Some of our competitors possess greater financial, marketing and research and development resources than we do.

Regulatory and Legal Matters

Regulatory Overview

Our operations are subject to extensive governmental regulation by virtually every country in which we operate including, most notably, in the U.S., at the federal, state and local levels. Although these regulations differ from country to country, in general, non-U.S. regulations are designed to accomplish the same objectives as U.S. regulations regarding the operation of dialysis clinics, laboratories and manufacturing facilities, the provision of quality health care for patients, the maintenance of occupational, health, safety and environmental standards and the provision of accurate reporting and billing for governmental payments and/or reimbursement. In the U.S., some states restrict ownership of health care providers by certain multi-level for-profit corporate groups or establish other regulatory barriers to the establishment of new dialysis clinics. Outside the U.S., each country has its own payment and reimbursement rules and procedures, and some countries prohibit ownership of health care providers or establish other regulatory barriers to direct ownership by foreign companies. In all jurisdictions, we work within the framework of applicable laws to establish alternative contractual arrangements to provide services to those facilities.

Any of the following matters could have a material adverse effect on our business, financial condition and results of operations:

failure to receive required licenses, certifications or other approvals for new facilities or significant delays in such receipt;

loss of various federal certifications or termination of licenses under the laws of any state or other governmental authority; and

changes resulting from health care reform or other government actions that reduce reimbursement or reduce or eliminate coverage for particular services we provide.

We must comply with all U.S., German and other legal and regulatory requirements under which we operate, including the U.S. federal Medicare and Medicaid Fraud and Abuse Amendments of 1977, as amended, generally referred to as the anti-kickback statute, the federal False Claims Act, the federal restrictions on certain physician referrals, commonly known as the Stark Law, U.S. federal rules under the Health Insurance Portability and Accountability Act of 1996 that protect the privacy of patient medical records and prohibit inducements to patients to select a particular health care provider (commonly known as HIPAA) and other fraud and abuse laws and similar state statutes, as well as similar laws in other countries. Moreover, there can be

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no assurance that applicable laws, or the regulations thereunder, will not be amended, or that enforcement agencies or the courts will not make interpretations inconsistent with our own, any one of which could have a material adverse effect on our business, reputation, financial condition and results. Sanctions for violations of these statutes may include criminal or civil penalties, such as imprisonment, fines or forfeitures, denial of payments, and suspension or exclusion from the Medicare and Medicaid programs. In the U.S., these laws have been broadly interpreted by a number of courts, and significant government funds and personnel have been devoted to their enforcement because such enforcement has become a high priority for the federal government and some states. Our company, and the health care industry in general, will continue to be subject to extensive federal, state and foreign regulation, the full scope of which cannot be predicted.

Fresenius Medical Care Holdings has entered into a corporate integrity agreement with the U.S. government, which requires that Fresenius Medical Care Holdings staff and maintain a comprehensive compliance program, including a written code of conduct, training programs and compliance policies and procedures. The corporate integrity agreement requires annual audits by an independent review organization and periodic reporting to the government. The corporate integrity agreement permits the U.S. government to exclude Fresenius Medical Care Holdings and its subsidiaries from participation in U.S. federal health care programs and impose fines if there is a material breach of the agreement that is not cured by Fresenius Medical Care Holdings within thirty days after Fresenius Medical Care Holdings receives written notice of the breach.

Product Regulation

U.S.

In the U.S., the Food and Drug Administration (FDA) and comparable state regulatory agencies impose requirements on certain of our subsidiaries as a manufacturer and a seller of medical products and supplies under their jurisdiction. These require that products be manufactured in accordance with Good Manufacturing Practices and that we comply with FDA requirements regarding the design, safety, advertising, labeling, recordkeeping distribution, and reporting of adverse events related to the use of our products. In addition, in order to clinically test, produce and market certain medical products and other disposables (including hemodialysis and peritoneal dialysis equipment and solutions, dialyzers, bloodlines and other disposables) for human use, we must satisfy mandatory procedures and safety and efficacy requirements established by the FDA or comparable state and foreign governmental agencies. Such rules generally require that products be approved by the FDA as safe and effective for their intended use prior to being marketed. Our peritoneal dialysis solutions have been designated as drugs by the FDA and, as such, are subject to additional FDA regulation under the Food, Drug and Cosmetic Act of 1938.

Germany and Other Non-U.S.

Most countries maintain different regulatory regimes for pharmaceutical products and for medical devices. In each regime, there are regulations governing manufacturers and distributors, as well as regulations governing the final products manufactured and distributed. Treaties or other international law and standards and guidelines under treaties or laws may supplement or supersede individual country regulations.

Some of our products, such as peritoneal dialysis solutions, are considered pharmaceuticals. The European Union has issued a directive on pharmaceuticals, No. 65/65/ EWG (January 26, 1965), as amended. Each member of the European Union is responsible for conforming its law to comply with this directive. In Germany the German Drug Law (*Arzneimittelgesetz*) which implements European Union requirements, is the primary regulation applicable to pharmaceutical products.

The provisions of the German Drug Law are typical of the legal standards in other European countries. The German Drug Law states the requirements for the authorization of a company to manufacture pharmaceuticals. A manufacturer must, among other requirements, appoint pharmacists or physicians to be responsible for the quality, safety and efficacy of the pharmaceuticals. At least five responsible persons must be appointed: a marketing manager, a quality control manager, a manufacturing manager, a safety officer, and a drug information

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officer. Each of these persons may be held personally liable under German criminal laws for violations of the German Drug Law.

International guidelines also govern the manufacture of pharmaceuticals and, in many cases, overlap with national requirements. In particular, the Pharmaceutical Inspection Convention, an international treaty, contains rules which are binding on most countries in which pharmaceuticals are manufactured. Among other things, the Pharmaceutical Inspection Convention establishes requirements for Good Manufacturing Practices which are then adopted at the national level. Another international guideline, which is non-binding, is the ISO 9000-9004 system for assuring quality control. This system is more detailed than Good Manufacturing Practices. Compliance entitles the manufacturer to utilize the CE certification of quality control. In addition to regulating the manufacture of pharmaceuticals, countries directly regulate marketing of the pharmaceuticals produced. A drug needs to be registered and authorized in every country in which it is distributed. European Union rules govern the conditions for a registration, such as pre-clinical and clinical testing.

Historically, medical devices have not been regulated as strictly as pharmaceuticals, but more stringent regulatory schemes have been adopted during the last decade. In 1995, Germany implemented the European Union's Medical Devices Directive when it adopted the Medical Devices Act (*Medizinproduktegesetz*), which is similar in many ways to the German Drug Law. This Directive applies to both the manufacturer's quality control system and the products' technical design. Depending on the class of medical devices, a manufacturer may choose alternative regulatory modules to demonstrate compliance with European Union provisions. To assure and demonstrate the high quality standards and performance of our operations, we have subjected our entire European business to the most comprehensive procedural module, which is also the fastest way to launch a new product in the European Union. This module requires the certification of a full quality management system by a notified body charged with supervising the quality management system. A notified body is a group accredited and monitored by governmental agencies that inspects manufacturing facilities and quality control systems at regular intervals and is authorized to carry out unannounced inspections.

When a company receives a European Union certificate for the quality management system of a particular facility, it may assess whether products developed and manufactured in the facility satisfy European Union requirements. European Union requirements for products are laid down in harmonized European Union standards and include conformity to safety requirements, physical and biological properties, construction and environmental properties, and information supplied by the manufacturer. A manufacturer must demonstrate conformity to these requirements by pre-clinical tests, biocompatibility tests, qualification of products and packaging, risk analysis and well-conducted clinical investigations approved by ethics committees.

A manufacturer having a European Union-certified full quality management system has to declare and document conformity of its products to the harmonized European standards. If able to do so, the manufacturer must put a "CE" mark on the products. The CE mark, which stands for *Conformité Européenne*, demonstrates compliance with the relevant European Union requirements. Products subject to these provisions that do not bear the "CE" mark cannot be imported, sold or distributed within the European Union.

Our Series 4008, 4008B, 4008E dialysis machines and their therapy modifications, our PD-NIGHT cyler, and our other medical devices distributed in the European market, as well as our dialysis filters and dialysis tubing systems and accessories, all bear the "CE" mark. We expect to continue to obtain additional certificates as they are required.

Facilities and Operational Regulation

U.S.

The Clinical Laboratory Improvement Amendments of 1988 (CLIA) subjects virtually all clinical laboratory testing facilities, including ours, to the jurisdiction of the Department of Health and Human Services. CLIA establishes national standards for assuring the quality of laboratories based upon the complexity of testing performed by a laboratory. Certain of our operations are also subject to federal laws governing the repackaging and dispensing of drugs and the maintenance and tracking of certain life sustaining and life-supporting equipment.

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Our operations are subject to various U.S. Department of Transportation, Nuclear Regulatory Commission and Environmental Protection Agency requirements and other federal, state and local hazardous and medical waste disposal laws. As currently in effect, laws governing the disposal of hazardous waste do not classify most of the waste produced in connection with the provision of dialysis, or laboratory services as hazardous, although disposal of nonhazardous medical waste is subject to specific state regulation. Our operations are also subject to various air emission and wastewater discharge regulations.

Federal, state and local regulations require us to meet various standards relating to, among other things, the management of facilities, personnel qualifications and licensing, maintenance of proper records, equipment, quality assurance programs, the operation of pharmacies, and dispensing of controlled substances. All of our operations in the U.S. are subject to periodic inspection by federal and state agencies and other governmental authorities to determine if the operations, premises, equipment, personnel and patient care meet applicable standards. To receive Medicare reimbursement, our dialysis centers, renal diagnostic support business and laboratories must be certified by CMS. All of our dialysis centers, and laboratories that furnish Medicare services have the required certification.

Certain of our facilities and certain of their employees are also subject to state licensing statutes and regulations. These statutes and regulations are in addition to federal and state rules and standards that must be met to qualify for payments under Medicare, Medicaid and other government reimbursement programs. Licenses and approvals to operate these centers and conduct certain professional activities are customarily subject to periodic renewal and to revocation upon failure to comply with the conditions under which they were granted.

Occupational Safety and Health Administration (OSHA) regulations require employers to provide employees who work with blood or other potentially infectious materials with prescribed protections against blood-borne and air-borne pathogens. The regulatory requirements apply to all health care facilities, including dialysis centers and laboratories, and require employers to make a determination as to which employees may be exposed to blood or other potentially infectious materials and to have in effect a written exposure control plan. In addition, employers are required to provide hepatitis B vaccinations, personal protective equipment, blood-borne pathogens training, post-exposure evaluation and follow-up, waste disposal techniques and procedures, engineering and work practice controls and other OSHA-mandated programs for blood-borne and air-borne pathogens.

Some states in which we operate have certificate of need (CON) laws that require any person or entity seeking to establish a new health care service or to expand an existing service to apply for and receive an administrative determination that the service is needed. We currently operate in 13 states, as well as the District of Columbia and Puerto Rico that have CON laws applicable to dialysis centers. These requirements could, as a result of a state s internal determination of its dialysis services needs, prevent entry to new companies seeking to provide services in these states, and could constrain our ability to expand our operations in these states.

Germany and Other Non-U.S.

Countries outside of the U.S. possess a wide variety of operational regulation at disparate levels. Accordingly, our operations are subject to very different regulations in different countries. Most countries regulate dialysis clinic operating conditions and product manufacturing.

We are subject to a broad spectrum of regulation. Our operations must comply with various environmental and transportation regulations in the various countries in which we operate. Our manufacturing facilities and dialysis clinics are also subject to various standards relating to, among other things, facilities, management, personnel qualifications and licensing, maintenance of proper records, equipment, quality assurance programs, the operation of pharmacies, the protection of workers from blood-borne diseases and the dispensing of controlled substances. All of our operations are subject to periodic inspection by various governmental authorities to determine if the operations, premises, equipment, personnel and patient care meet applicable standards. Our dialysis clinic operations and our related activities generally require licenses, which are subject to periodic renewal and may be revoked for violation of applicable regulatory requirements.

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In addition, many countries impose various investment restrictions on foreign companies. For instance, government approval may be required to enter into a joint venture with a local partner. Some countries do not permit foreign investors to own a majority interest in local companies or require that companies organized under their laws have at least one local shareholder. Investment restrictions therefore affect the corporate structure, operating procedures and other characteristics of our subsidiaries and joint ventures in these and other countries.

We believe our facilities are currently in compliance in all material respects with the applicable national and local requirements in the jurisdictions in which they operate.

Reimbursement

U.S.

Dialysis Services. Our dialysis centers provide outpatient hemodialysis treatment and related services for ESRD patients. In addition, some of the Company's centers offer services for the provision of peritoneal dialysis and hemodialysis treatment at home, and dialysis for hospitalized patients.

The Medicare program is the primary source of Dialysis Services revenues from dialysis treatment. For example, in 2003, approximately 55% of Dialysis Services revenues resulted from Medicare's ESRD program. As described below, Dialysis Services is reimbursed by the Medicare program in accordance with the Composite Rate for certain products and services rendered at our dialysis centers. As described hereinafter, other payment methodologies apply to Medicare reimbursement for other products and services provided at our dialysis centers and for products (such as those sold by us) and support services furnished to ESRD patients receiving dialysis treatment at home (such as those of Dialysis Products). Medicare reimbursement rates are fixed in advance and are subject to adjustment from time to time by the U.S. Congress. Although this form of reimbursement limits the allowable charge per treatment, it provides us with predictable per treatment revenues.

Certain items and services that we furnish at our dialysis centers are not included in the Composite Rate and are eligible for separate Medicare reimbursement, typically on the basis of established fee schedule amounts. Such items and services include certain drugs (such as EPO), blood transfusions and certain diagnostic tests.

Medicare payments are subject to change by legislation, regulations and pursuant to deficit reduction measures. The Composite Rate was unchanged from commencement of the ESRD program in 1972 until 1983. From 1983 through December 1990, numerous congressional actions resulted in a net reduction of the average reimbursement rate from \$138 per treatment in 1983 to approximately \$125 per treatment in 1990. Congress increased the ESRD reimbursement rate, effective January 1, 1991, to an average rate of \$126 per treatment. Effective January 1, 2000, the reimbursement rate was increased by 1.2%. In December 2000 an additional increase of 2.4% was approved for the year 2001. Accordingly, there was a 1.2% reimbursement increase on January 1, 2001. A second increase was delayed until April 1, 2001, when rates were increased 1.6% to make up for the delay.

On December 8, 2003, the Medicare Prescription Drug, Modernization and Improvement Act of 2003 was enacted (the Medicare Modernization Act). This law makes several modifications affecting payment for dialysis services. First, it will increase the composite rate for renal dialysis facilities by 1.6% on January 1, 2005. Second, it requires the Secretary of the Department of Health and Human Services (the Secretary) to establish a new case-mix adjusted prospective payment system for dialysis services furnished on or after January 1, 2005. This system will adjust for a limited number of patient characteristics (the case-mix) and will include two components: (1) those services that currently comprise the Composite Rate; and (2) the difference between the Medicare payment rate for separately billable drugs and biologicals and the acquisition costs of those drugs and biologicals, as determined by OIG reports. The Secretary is required to adjust the basic case-mix adjusted system payment rates by a geographic index. Separate payment would continue to be made for drugs and biologicals that currently are billed separately. All separately billable drugs and biologicals will be reimbursed based upon acquisition cost in 2005. Beginning in 2006, the Secretary is authorized to set payment for all separately billed drugs and biologicals at either acquisition cost or average sales price. Third, the Secretary is required to establish a three-year demonstration project to test the use of a fully case-mix adjusted payment system for ESRD services, beginning January 1, 2006. Under this project, separately billable drugs and biologicals and related clinical

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laboratory tests would be bundled into the facility composite rate. Participating facilities would receive an additional 1.6% composite rate increase.

We are unable to predict what, if any, future changes may occur in the rate of Medicare reimbursement. Any significant decreases in the Medicare reimbursement rates could have a material adverse effect on our provider business and, because the demand for products is affected by Medicare reimbursement, on our products business. Increases in operating costs that are affected by inflation, such as labor and supply costs, without a compensating increase in reimbursement rates, also may adversely affect our business and results of operations.

For Medicare-primary patients, Medicare is responsible for payment of 80% of the Composite Rate set by CMS for dialysis treatments and the patient or third-party insurance payors, including employer-sponsored health insurance plans, commercial insurance carriers and the Medicaid program, are responsible for paying any co-payment amounts for approved services not paid by Medicare (typically the annual deductible and 20% co-insurance), subject to the specific coverage policies of such payors. Each third-party payor, including Medicaid, makes payment under contractual or regulatory reimbursement provisions which may or may not cover the full 20% co-payment or annual deductible. Where the patient has no third-party insurance or the third party insurance does not cover the co-payment or deductible, the patient is responsible for paying the co-payments or the deductible, which we frequently do not fully collect despite reasonable collection efforts. Under an advisory opinion from the Office of the Inspector General, subject to specified conditions, we and other similarly situated providers may make contributions to a non-profit organization that has agreed to make premium payments for supplemental medical insurance and/ or medigap insurance on behalf of indigent ESRD patients, including some of our patients.

Laboratory Tests. Spectra Renal Management obtains a substantial portion of its net revenue from Medicare, which pays for clinical laboratory services provided to dialysis patients in two ways.

First, payment for certain routine tests is included in the Composite Rate paid to our dialysis centers. As to such services, the dialysis centers obtain the services from a laboratory and pay the laboratory for such services. In accordance with industry practice, Spectra Renal Management usually provides such testing services under capitation agreements with its customers pursuant to which it bills a fixed amount per patient per month to cover the laboratory tests included in the Composite Rate at the designated frequencies. In October 1994, the Office of Inspector General issued a special fraud alert in which it stated its view that the industry practice of providing tests covered by the Composite Rate at below fair market value raised issues under the anti-kickback statutes, as such an arrangement with an ESRD facility appeared to be an offer of something of value (Composite Rate tests at below market value) in return for the ordering of additional tests billed directly to Medicare. See Anti-kickback Statutes, False Claims Act, Stark Law and Fraud and Abuse Laws for a description of this statute.

Second, laboratory tests performed by Spectra Renal Management for Medicare beneficiaries that are not included in the Composite Rate are separately billable directly to Medicare. Such tests are paid at 100% of the Medicare fee schedule amounts, which are limited by national ceilings on payment rates, called National Limitation Amounts (NLAs). Congress has periodically reduced the fee schedule rates and the NLAs, with the most recent reductions in the NLAs occurring in January 1998. (As part of the Balanced Budget Act of 1997, Congress lowered the NLAs from 76% to 74% effective January 1, 1998.) Congress has also approved a five year freeze on the inflation updates based on the Consumer Price Index (CPI) for 1998-2002.

Erythropoetin (EPO). Any of the following could adversely affect our business, and results of operations, possibly materially:

future changes in the EPO reimbursement rate without offsetting changes to the Medicare composite rate;

inclusion of EPO in the Medicare composite rate without offsetting increases to such rate;

changes in the typical dosage per administration;

increases in the cost of EPO after our current supply contract expires; or

reduction by the manufacturer of EPO of the amount of overfill in the EPO vials.

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Coordination of Benefits. Medicare entitlement begins for most patients in the fourth month after the initiation of chronic dialysis treatment at a dialysis center. During the first three months, considered to be a waiting period, the patient or patient's insurance, Medicaid or a state renal program are responsible for payment.

Patients who are covered by Medicare and are also covered by an employer group health plan (EGHP) are subject to a 30-month coordination period during which the EGHP is the primary payor and Medicare the secondary payor. During this coordination period the EGHP pays a negotiated rate or in the absence of such a rate, our standard rate or a rate defined by its plan documents. The EGHP payments are generally higher than the Medicare Composite Rate. EGHP insurance, when available, will therefore generally cover as the primary payor a total of 33 months, the 3-month waiting period plus the 30-month coordination period.

Possible Changes in Medicare. Legislation or regulations may be enacted in the future that could substantially modify or reduce the amounts paid for services and products offered by us and our subsidiaries. It is also possible that statutes may be adopted or regulations may be promulgated in the future that impose additional eligibility requirements for participation in the federal and state health care programs. Such new legislation or regulations may adversely affect our businesses and results of operations, possibly materially.

Germany and Other Non-U.S.

As a global company delivering dialysis care and dialysis products in more than 100 countries world wide, we face the challenge of addressing the needs of dialysis patients in widely varying economic and health care environments.

Health care systems and reimbursement structures for ESRD treatment vary by country. In general, the government pays for health care and finances its payments through taxes and other sources of government income, from social contributions, or a combination of those sources. However, not all health care systems provide for dialysis treatment. In many developing countries, only limited subsidies from government or charitable institutions are available, and dialysis patients must finance all or substantially all of the cost of their treatment. In some countries patients in need of dialysis do not receive treatment on a regular basis but rather when the financial resources allow it.

In the major European and British Commonwealth countries, health care systems are generally based on one of two models. The German model is based on mandatory employer and employee contributions dedicated to health care financing. The British model provides a national health care system funded by taxes, with the result that funds allocated to health care may vary from year to year. Within these systems provision for the treatment of dialysis has been made either through allocation of a national budget or a billing system reimbursing on a fee-for-service basis. The health care systems of Japan, France, Switzerland and the Netherlands are based on the German model. Canada, Scandinavia, Italy and Spain established their national health services using the British model.

Ownership of health care providers and, more specifically dialysis care providers, varies within the different systems and from country-to-country. In Europe almost 60% of the clinics providing dialysis care and services are publicly owned, more than 30% are privately owned and approximately 10% belong to a health care organization. It should be noted that health care organizations treating a significant patient population operate only in Germany, France and Spain. Publicly operated clinics care for almost 100% of the dialysis populations in Canada and Australia. Within Europe, nearly 100% of the dialysis population is treated in public clinics in the United Kingdom, the Netherlands and the Scandinavian countries, while more than 50% of dialysis clinics are privately owned in Spain and Portugal.

In Latin America privately owned clinics predominate, constituting more than 60% of all clinics providing dialysis care while in Asia, with the exception of Japan, publicly owned clinics are predominant. As in the U.S., only approximately 15% of dialysis clinics in Japan are publicly operated. Unlike the U.S., however, Japan has a premium-based, mandatory social insurance system, and the structure of its health care system is more closely comparable to the German system.

Financing policies for ESRD treatment also differ from country-to-country. In countries with a health care system that includes provisions for ESRD patient care, treatment is generally financed through a government

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budget allocation or on a fee-for-service basis. A few European countries have introduced payment systems based on fixed fees charged according to the disease related group, an arrangement similar to capitation. This basis for payment was adopted from the United States where it was implemented as a method to curtail costs.

Treatment components included in the cost of dialysis may vary from country-to-country, depending on the structure and cost allocation principles. Where treatment is reimbursed on a fee-for-service basis, reimbursement rates are allocated in accordance with the type of treatment performed. We believe that it is not appropriate to calculate a global reimbursement amount, because the services and costs for which reimbursement is provided in any such global amount would be likely to bear little relation to the actual reimbursement system in any one country. Generally, in countries with established dialysis programs, reimbursements range from \$100 to more than \$300 per treatment. However, a comparison from country to country would not be meaningful if made in the absence of a detailed analysis of the cost components reimbursed, services rendered and the structure of the dialysis clinic in each country being compared.

Health care expenditures are consuming an ever increasing portion of gross domestic product worldwide. In the developed economies of Europe, Asia and Latin America, health care spending is in the range of 5%-14% of gross domestic product. As in the U.S., dialysis costs consume a disproportionately high amount of health care spending and these costs may be considered a target for implementation of cost containment measures. Today, there is increasing awareness of the correlation between the quality of care delivered in the dialysis unit and the total health care expenses incurred by the dialysis patient. Accordingly, developments in reimbursement policies include higher reimbursement rates for practices which are believed to improve the overall state of health of the ESRD patient and reduce the need for additional medical treatment.

Anti-kickback Statutes, False Claims Act, Health Care Fraud, Stark Law and Fraud and Abuse Laws in North America

Some of our operations are subject to federal and state statutes and regulations governing financial relationships between health care providers and potential referral sources and reimbursement for services and items provided to Medicare and Medicaid patients. Such laws include the anti-kickback statute, health care fraud statutes, the False Claims Act, the Stark Law, other federal fraud and abuse laws and similar state laws. These laws apply because our Medical Directors and other physicians with whom we have financial relationships refer patients to, and order diagnostic and therapeutic services from, our dialysis centers and other operations. As is generally true in the dialysis industry, at each dialysis facility a small number of physicians account for all or a significant portion of the patient referral base. An ESRD patient generally seeks treatment at a center that is convenient to the patient and at which the patient's nephrologist has staff privileges.

Anti-kickback Statutes

The federal anti-kickback statute establishes criminal prohibitions against and civil penalties for the knowing and willful solicitation, receipt, offer or payment of any remuneration, whether direct or indirect, in return for or to induce the referral of patients or the ordering or purchasing of items or services payable in whole or in part under Medicare, Medicaid or other federal health care programs. Sanctions for violations of the anti-kickback statute include criminal and civil penalties, such as imprisonment or criminal fines of up to \$25,000 per violation, and civil penalties of up to \$50,000 per violation, and exclusion from the Medicare or Medicaid programs and other federal programs. In addition, certain provisions of federal criminal law that may be applicable provide that if a corporation is found guilty of a criminal offense it may be fined no more than twice any pecuniary gain to the corporation, or, in the alternative, no more than \$500,000 per offense.

Some states also have enacted statutes similar to the anti-kickback statute, which may include criminal penalties, applicable to referrals of patients regardless of payor source, and may contain exceptions different from state to state and from those contained in the federal anti-kickback statute.

False Claims Act and Related Criminal Provisions

The federal False Claims Act (the False Claims Act) imposes civil penalties for knowingly making or causing to be made false claims with respect to governmental programs, such as Medicare and Medicaid, for

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services billed but not rendered, or for misrepresenting actual services rendered, in order to obtain higher reimbursement. Moreover, private individuals may bring qui tam or whistle blower suits against providers under the False Claims Act, which authorizes the payment of a portion of any recovery to the individual bringing suit. Such actions are initially required to be filed under seal pending their review by the Department of Justice. A few federal district courts have recently interpreted the False Claims Act as applying to claims for reimbursement that violate the anti-kickback statute or federal physician self-referral law under certain circumstances. The False Claims Act generally provides for the imposition of civil penalties of \$5,500 to \$11,000 per claim and for treble damages, resulting in the possibility of substantial financial penalties for small billing errors that are replicated in a large number of claims, as each individual claim could be deemed to be a separate violation of the False Claims Act. Criminal provisions that are similar to the False Claims Act provide that if a corporation is convicted of presenting a claim or making a statement that it knows to be false, fictitious or fraudulent to any federal agency it may be fined not more than twice any pecuniary gain to the corporation, or, in the alternative, no more than \$500,000 per offense. Some states also have enacted statutes similar to the False Claims Act which may include criminal penalties, substantial fines, and treble damages.

The Health Insurance Portability and Accountability Act of 1996

HIPAA was enacted in August 1996 and expanded federal fraud and abuse laws by increasing their reach to all federal health care programs, establishing new bases for exclusions and mandating minimum exclusion terms, creating an additional exception to the anti-kickback penalties for risk-sharing arrangements, requiring the Secretary of Health and Human Services to issue advisory opinions, increasing civil money penalties to \$10,000 (formerly \$2,000) per item or service and assessments to three times (formerly twice) the amount claimed, creating a specific health care fraud offense and related health fraud crimes, and expanding investigative authority and sanctions applicable to health care fraud. It also prohibits a provider from offering anything of value which the provider knows or should know would be likely to induce the patient to select the provider.

The law expands criminal sanctions for health care fraud involving any governmental or private health benefit program, including freezing of assets and forfeiture of property traceable to commission of a health care offense.

HIPAA included a health care fraud provision which prohibits knowingly and willfully executing a scheme or artifice to defraud any health care benefit program, which includes any public or private plan or contract affecting commerce under which any medical benefit, item, or service is provided to any individual, and includes any individual or entity who is providing a medical benefit, item, or service for which payment may be made under the plan or contract.

HIPAA regulations establish national standards for certain electronic health care transactions, the use and disclosure of certain individually identifiable patient health information, and the security of the electronic systems maintaining this information. These are commonly known as the HIPAA transaction and code set standards, privacy standards, and security standards. Health insurance payers and healthcare providers like us must comply with the new HIPAA standards. Violations of these HIPAA standards may include civil money penalties and potential criminal sanctions.

Balanced Budget Act of 1997

The Balanced Budget Act of 1997 (the BBA) contained material adjustments to both the Medicare and Medicaid programs, as well as further expansion of the federal fraud and abuse laws. Specifically, the BBA created a civil monetary penalty for violations of the federal anti-kickback statute whereby violations will result in damages equal to three times the amount involved as well as a penalty of \$50,000 per violation. In addition, the new provisions expanded the exclusion requirements so that any person or entity convicted of three health care offenses is automatically excluded from federally funded health care programs for life. Individuals or entities convicted of two offenses are subject to mandatory exclusion of 10 years, while any provider or supplier convicted of any felony may be denied entry into the Medicare program by the Secretary of HHS if deemed to be detrimental to the best interests of the Medicare program or its beneficiaries.

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The BBA also provides that any person or entity that arranges or contracts with an individual or entity that has been excluded from a federally funded health care program will be subject to civil monetary penalties if the individual or entity knows or should have known of the sanction.

Stark Law

The original Stark Law, known as Stark I and enacted as part of the Omnibus Budget Reconciliation Act (OBRA) of 1989, prohibits a physician from referring Medicare patients for clinical laboratory services to entities with which the physician (or an immediate family member) has a financial relationship, unless an exception applies. Sanctions for violations of the Stark Law may include denial of payment, refund obligations, civil monetary penalties and exclusion of the provider from the Medicare and Medicaid programs. The Stark Law prohibits the entity receiving the referral from filing a claim or billing for services arising out of the prohibited referral.

Provisions of OBRA 93, known as Stark II, amended Stark I to revise and expand upon various statutory exceptions, to expand the services regulated by the statute to a list of Designated Health Services, and expanded the reach of the statute to the Medicaid program. The provisions of Stark II generally became effective on January 1, 1995, with the first phase of Stark II regulations finalized on January 4, 2001. Most portions of the first phase regulations became effective in 2002. The additional Designated Health Services include: physical therapy services; occupational therapy services; radiology services and certain other imaging services; durable medical equipment and supplies; parenteral and enteral nutrients, equipment and supplies; prosthetics, orthotics, and prosthetic devices and supplies; home health services; outpatient prescription drugs; and inpatient and outpatient hospital services. The first phase of the final regulations implementing the Stark Law contains an exception for EPO and certain other dialysis-related outpatient prescription drugs furnished in or by an ESRD facility under many circumstances. Further, the final regulations also adopt a definition of durable medical equipment which effectively excludes ESRD equipment and supplies from the category of Designated Health Services. Phase II of the final regulations to the Stark Law, which will address many of the compensation exceptions, has not yet been released.

Several states in which we operate have enacted self-referral statutes similar to the Stark Law. Such state self-referral laws may apply to referrals of patients regardless of payor source and may contain exceptions different from each other and from those contained in the Stark Law.

Other Fraud and Abuse Laws

Our operations are also subject to a variety of other federal and state fraud and abuse laws, principally designed to ensure that claims for payment to be made with public funds are complete, accurate and fully comply with all applicable program rules.

The civil monetary penalty provisions are triggered by violations of numerous rules under the Medicare statute, including the filing of a false or fraudulent claim and billing in excess of the amount permitted to be charged for a particular item or service. Violations may also result in suspension of payments, exclusion from the Medicare and Medicaid programs, as well as other federal health care benefit programs, or forfeiture of assets.

In addition to the statutes described above, other criminal statutes may be applicable to conduct that is found to violate any of the statutes described above.

Health Care Reform

Health care reform is considered by many countries to be a national priority. In the U.S., members of Congress from both parties and officials from the executive branch are continuing to consider many health care proposals, some of which are comprehensive and far-reaching in nature. Several states are also currently considering health care proposals. We cannot predict what additional action, if any, the federal government or any state may ultimately take with respect to health care reform or when any such action will be taken. Health care reform may bring radical changes in the financing and regulation of the health care industry, which could have a material adverse effect on our business and the results of our operations.

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C. Organizational Structure

The following chart shows our organizational structure and our significant subsidiaries. Fresenius Medical Care Holdings, Inc. conducts its business as Fresenius Medical Care North America.

Table of Contents**D. Property, plant and equipment***Property*

The table below describes our principal facilities. We do not own the land and buildings comprising our principal facilities in Germany. Rather, we lease those facilities on a long-term basis from Fresenius AG or one of its affiliates. This lease is described under Item 7.B. Related Party Transactions Real Property Lease.

Location	Floor Area (Approximate Square Meters)	Currently Owned or Leased by Fresenius Medical Care	Lease Expiration	Use
Bad Homburg, Germany	5,374	leased	December 2006	Corporate headquarters and administration
St. Wendel, Germany	49,732	leased	December 2006	Manufacture of polysulfone membranes and dialyzers, bloodlines, and peritoneal dialysis solutions; research and development
Schweinfurt, Germany	15,717	leased	December 2006	Manufacture of hemodialysis machines and peritoneal dialysis cyclers; research and development
L. Arbresle, France	13,524	owned		Manufacture of polysulfone dialyzers and special filters dry hemodialysis concentrates
Palazzo Pignano, Italy	66,550	owned		Manufacture of bloodlines and tubing
Nottinghamshire, United Kingdom	5,110	owned		Manufacture of hemodialysis concentrate solutions
Barcelona, Spain	2,000	owned		Manufacture of peritoneal dialysis bags and concentrates
Ankara, Turkey	1,000	leased	February 2009	Manufacture of liquid hemodialysis concentrate solutions
Tunisia	491	leased	December 2020	Manufacture of liquid hemodialysis concentrate solutions
Buenos Aires, Argentina	10,100	owned		Manufacture of hemodialysis concentrate solutions
Rio de Janeiro, Brazil	1,159	leased	Month to Month	Manufacture of hemodialysis concentrate solutions
Oita, Japan ⁽¹⁾	24,083	owned		Manufacture of polysulfone membranes, dialyzers, dialysis solutions, dialysis machine components and assembly of dialysis machines
Hong Kong	1,013	Leased	February 2006	Corporate headquarter and administration
Milson Point, Australia	557	leased	November 2007	Administration
Smithfield, Australia	5,350	owned		Manufacture of hemodialysis concentrate solutions Warehouse
Pusat, KL, Malaysia	4,060	leased	November 2004	Administration
Taipei, Taiwan	5,940	leased	December 2006	Administration
Tokyo, Japan	1,153	leased	December 2006 with 3-year renewal option	Administration
Inukai, Japan	2,142	owned		Manufacture of polysulfone dialyzers and filters
Buzen, Japan	8,369	owned		

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Location	Floor Area (Approximate Square Meters)	Currently Owned or Leased by Fresenius Medical Care	Lease Expiration	Use
Seoul, South Korea	2,426	leased	March and August 2004	Administration
Lexington, Massachusetts	18,581	leased	October 2007 with 5-year renewal option	Corporate headquarters and administration North America
Walnut Creek, California	9,522	leased	June 2012 with 5-year renewal option	Manufacture of Hemodialysis machines and peritoneal dialysis cyclers; research and development; warehouse space
Ogden, Utah	41,807	owned		Manufacture polysulfone membranes and dialyzers and peritoneal dialysis solutions; research and development
Delran, New Jersey	3,902	leased	October 2004 with 5-year renewal option	Manufacture of liquid hemodialysis concentrate solutions
Perrysburg, Ohio	3,252	leased	August 2008	Manufacture of dry hemodialysis concentrates
Livingston, California	2,973	leased	October 2006 with a 5-year renewal option	Manufacture of liquid hemodialysis concentrates
Freemont, California	6,688	leased	May 2007 with 2-year renewal option	Clinical laboratory testing
Rockleigh, New Jersey	7,897	leased	June 2007 with two 5-year renewal options	Clinical laboratory testing
Miami, Florida	400	leased	Month to Month	Administration
Irving, Texas	6,503	leased	December 2010	Manufacture of liquid hemodialysis solution
Reynosa, Mexico	13,936	leased	June 2013	Manufacture of bloodlines

(1) We own 70% of the joint venture that owns this facility.

We lease most of our dialysis clinics, manufacturing, laboratory, warehousing and distribution and administrative and sales facilities in the U.S. and foreign countries on terms which we believe are customary in the industry. We own those dialysis clinics and manufacturing facilities that we do not lease.

For information regarding plans to expand our facilities and related capital expenditures, see Item 4.A. History and Development of the Company Capital Expenditures.

Item 5. Operating and Financial Review and Prospects

You should read the following discussion and analysis of the results of operations of Fresenius Medical Care AG and its subsidiaries in conjunction with our historical consolidated financial statements and related notes contained elsewhere in this report. Some of the statements contained below, including those concerning future revenue, costs and capital expenditures and possible changes in our industry and competitive and financial conditions include forward-looking statements. Because such statements involve risks and uncertainties, actual results may differ materially from the results which the forward looking statements express or imply.

Critical Accounting Policies

The Company's reported financial condition and results of operations are sensitive to accounting methods, assumptions and estimates that are the basis for our financial statements. The critical accounting policies, the judgments made in the creation and application of these policies, and the sensitivities of reported results to changes in accounting policies, assumptions and estimates are factors to be considered along with the Company's financial statements, and the discussion in Operating Results.

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Recoverability of Goodwill and Intangible Assets

The growth of our business through acquisitions has created a significant amount of intangible assets, including goodwill, trade names and management contracts. At December 31, 2003, the carrying amount of goodwill amounted to \$3,288 million and non-amortizable intangible assets amounted to \$431 million representing in total approximately 50% of our total assets.

In accordance with Statement of Financial Accounting Standards (SFAS) No. 142 *Goodwill and Other Intangible Assets* an annual impairment test of goodwill and non-amortizable intangible assets is performed at least once a year for each reporting unit, or if events occur or circumstances change that would indicate the carrying value might be impaired (See also Note 1g in our Consolidated Financial Statements).

To comply with the provisions of SFAS No. 142, the fair value of the reporting unit is compared to the reporting unit's carrying amount. We estimate the fair value of each reporting unit using estimated future cash flows for the unit discounted by a weighted average cost of capital specific to that unit. Estimated cash flows are based on our budgets for the next three years, and projections for the following years based on an expected growth rate. The growth rate is based on industry and internal projections. The discount rates reflect any inflation in local cash flows and risks inherent to each reporting unit.

If the fair value of the reporting unit is less than its carrying value, a second step is performed which compares the fair value of the reporting unit's goodwill to the carrying value of its goodwill. If the fair value of the goodwill is less than its carrying value, the difference is recorded as an impairment.

A prolonged downturn in the healthcare industry with lower than expected increases in reimbursement rates and/or higher than expected costs for providing healthcare services could adversely affect our estimated future cashflows. Future adverse changes in a reporting unit's economic environment could affect the discount rate. A decrease in our estimated future cash flows and/or a decline in the reporting units economic environment could result in impairment charges to goodwill and other intangible assets which could materially and adversely affect our future financial position and operating results.

Legal Contingencies

We are party to litigation relating to a number of matters as described in Note 20 *Legal Proceedings* in our Consolidated Financial Statements. The outcome of these matters may have a material effect on our financial position, results of operations or cash flows.

We regularly analyze current information including, as applicable, our defenses and provide accruals for probable contingent losses including the estimated legal expenses to resolve the matters. We use the resources of our internal legal department as well as external lawyers for the assessment. In making the decision, we consider the degree of probability of an unfavorable outcome and our ability to make a reasonable estimate of the amount of loss.

If an unfavorable outcome is probable but the amount of loss cannot be reasonably estimated by management, appropriate disclosure is provided, but no contingent losses are accrued. The filing of a suit or formal assertion of a claim or assessment does not automatically indicate that accrual of a loss may be appropriate.

Allowance for Doubtful Accounts

Trade accounts receivable are a significant asset of ours and the allowance for doubtful accounts is a significant estimate made by management. Trade accounts receivable were \$1,230 million and \$914 million at December 31, 2003 and 2002, respectively, net of allowances and after sales of accounts receivable under our accounts receivable facility described in Note 6 *Sale of Accounts Receivable* in our Consolidated Financial Statements. The allowance for doubtful accounts was \$166 million and \$160 million at December 31, 2003 and 2002, respectively. The majority of our receivables relates to our dialysis service business in North America.

Dialysis care revenues are recognized and billed at amounts estimated to be received under reimbursement arrangements with third party payors. Medicare and Medicaid programs are billed at pre-determined net

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realizable rates per treatment that are established by statute or regulation. Most non-governmental payors, including contracted managed care payors, are billed at the Company's standard rates for services net of contractual allowances to reflect the estimated amounts to be received under reimbursement arrangements with these payors.

Estimates for the allowances for doubtful accounts receivable from the dialysis service business are mainly based on past collection history. Specifically, the allowances for the North American operations are based on an analysis of collection experience, recognizing the differences between payors and aging of accounts receivable. From time to time, accounts receivable are reviewed for changes from the historic collection experience to ensure the appropriateness of the allowances. The allowances in the international segment and the products business are also based on estimates and consider various factors, including aging, creditor and past collection history.

A significant change in our collection experience, a deterioration in the aging of receivables and collection difficulties could require that we increase our estimate of the allowance for doubtful accounts. Any such additional bad debt charges could materially and adversely affect our future operating results.

Self Insurance Programs

The Company's largest subsidiary is partially self-insured for professional, product and general liability, auto and worker's compensation claims under which the company assumes responsibility for incurred claims up to predetermined amounts above which third party insurance applies. Reported balances for the year include estimates of the anticipated expense for claims incurred (both reported and incurred but not reported) based on historical experience and existing claim activity. This experience includes both the rate of claims incidence (number) and claim severity (cost) and is combined with individual claim expectations to estimate the reported amounts.

Financial Condition and Results of Operations

This section contains forward-looking statements. We made these forward-looking statements based on our management's expectations and beliefs concerning future events which may affect us, but we cannot assure that such events will occur or that the results will be as anticipated. Such statements include the matters that we described in the discussion in this report entitled "Forward-Looking Statements."

Our business is also subject to other risks and uncertainties that we describe from time to time in our public filings. Developments in any of these areas could cause our results to differ materially from the results that we or others have projected or may project.

Overview

We are engaged primarily in providing dialysis services and manufacturing and distributing products and equipment for the treatment of end-stage renal disease. In addition we perform clinical laboratory and renal diagnostic testing in the U.S. Dialysis is a lifesaving treatment for irreversible, lifelong end stage renal disease, and necessitates multiple treatments per week for the remainder of a patient's life. The provision of dialysis services and the distribution of dialysis products and equipment represents, based on our estimate, an over \$30 billion worldwide market and it is expected there will be annual patient growth of 5-7%. Patient growth is caused by factors such as the aging population, increasing incidence of diabetes and hypertension, improvements in treatment quality and improving standards of living in developing countries. Key to continued growth in revenue is our ability to attract new patients in order to increase the number of treatments performed each year. For that reason, we believe the number of treatments performed each year is a strong indicator of continued revenue growth and success. In addition the reimbursement environment significantly influences our business. In the past we experienced and also expect in the future generally stable reimbursements for dialysis services. This includes the balancing of unfavorable reimbursement changes in certain countries with favorable changes in other countries. The majority of treatments are paid by governmental institutions such as Medicare in the United States. As a consequence of the pressure to decrease health care costs, reimbursement rate increases have been limited. Our ability to influence the pricing of our services is limited. Profitability depends on our ability to manage rising labor, drug and supply costs.

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On December 8, 2003, the Medicare Prescription Drug, Modernization and Improvement Act of 2003 was enacted. Effective January 1, 2005, (1) the dialysis composite rate will increase 1.6%; (2) payments for separately billable dialysis-related medications will be based on acquisition cost, and an amount equal to the difference between acquisition cost and what would have been received under the 2003 reimbursement methodology will be added to the composite rate, and this add-back amount will be subject to an annual update based on the growth in drug spending; and (3) composite rate payments will be subject to a case mix adjustment system, resulting in higher composite rate payments for patients with more complicated medical cases. We expect the rate increase to have a positive effect on our results, and the two remaining changes to have no effect.

Our operations are geographically organized and accordingly we have identified three operating segments, North America, International, and Asia Pacific. For reporting purposes, we have aggregated the International and Asia Pacific segments as International. We aggregated these segments due to their similar economic characteristics. These characteristics include same services provided and same products sold, same type patient population, similar methods of distribution of products and services and similar economic environments. Our management board member responsible for the profitability and cash flow of each segment's various businesses supervises the management of each operating segment. The accounting policies of the operating segments are the same as those we apply in preparing our consolidated financial statements under accounting principles generally accepted in the United States (U.S. GAAP).

Our management evaluates each segment using a measure that reflects all of the segment's controllable revenues and expenses. Our management believes the most appropriate measure in this regard is operating income, referred to in previous filings as earnings before interest and taxes, or EBIT, which measures our source of earnings. Financing is a corporate function which segments do not control. Therefore, we do not include interest expense relating to financing as a segment measurement. We also regard income taxes to be outside the segments' control. In addition to operating income, our management also believes that earnings before interest, taxes, depreciation and amortization, or EBITDA, is helpful for investors as a measurement of our segments' ability to generate cash and to service our financing obligations. EBITDA is also the basis for determining compliance with certain covenants contained in our senior credit agreement and the indentures relating to our outstanding trust preferred securities. You should not consider segment EBITDA to be an alternative to net earnings determined in accordance with U.S. GAAP or to cash flow from operations, investing activities or financing activities. We believe that operating income is the GAAP financial measure most directly comparable to our computation of EBITDA by segment, and the information in the table below under Results of Operations reconciles EBITDA for each of our reporting segments to operating income calculated in accordance with U.S. GAAP. See also Note 23 of the Notes to Condensed Consolidated Financial Statements. Our discussions relating to our consolidated financial position and results of operations for 2001 reflect the effects of the special charge recorded in that year. **The discussion of the disaggregated results of operations of the North America segment and the discussion under Corporate exclude the effect of those special charges.**

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The following tables summarize our financial performance and certain operating results by reporting segment for the periods indicated. Inter-segment sales primarily reflect sales of medical equipment and supplies from the International segment to the North America segment. In order to facilitate a year-to-year comparison, goodwill adjusted figures for 2001, as if SFAS No. 142, *Goodwill and Other Intangible Assets*, had been adopted as of January 1, 2001, have also been provided. We prepared the information using a management approach, consistent with the basis and manner in which our management internally disaggregates financial information to assist in making internal operating decisions and evaluating management performance.

	For the years ended December 31,			
	2003	2002	2001 ^(a)	2001
	(in millions)			
Total revenue				
North America	\$3,857	\$3,750	\$3,604	\$3,604
International	1,709	1,363	1,281	1,281
	<u>5,566</u>	<u>5,113</u>	<u>4,885</u>	<u>4,885</u>
Totals				
North America	2	2	2	2
International	36	27	24	24
	<u>38</u>	<u>29</u>	<u>26</u>	<u>26</u>
Totals				
North America	3,855	3,748	3,602	3,602
International	1,673	1,336	1,257	1,257
	<u>5,528</u>	<u>5,084</u>	<u>4,859</u>	<u>4,859</u>
Totals				
North America	652	630	693	693
International	349	292	292	292
Special charge for legal matters			(258)	(258)
Corporate	(27)	(16)	(24)	(24)
	<u>974</u>	<u>906</u>	<u>703</u>	<u>703</u>
Totals				
North America	120	139	140	247
International	95	70	62	76
Corporate	2	2	1	1
	<u>217</u>	<u>211</u>	<u>203</u>	<u>324</u>
Totals				
North America	532	491	553	446
International	254	222	230	216
Special charge for legal matters			(258)	(258)
Corporate	(29)	(18)	(25)	(25)
	<u>757</u>	<u>695</u>	<u>500</u>	<u>379</u>
Totals				

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Interest income	19	18	14	14
Interest expense	(230)	(244)	(237)	(237)
Income tax expense	(213)	(175)	(109)	(91)
Minority interest	(2)	(4)	(2)	(2)
Net income	\$ 331	\$ 290	\$ 166	\$ 63

(a) Financial performance and certain operating results by principal business segment for the year ended December 31, 2001 as if SFAS No. 142, *Goodwill and Other Intangible Assets* was adopted on January 1, 2000. Management believes that presentation of our 2001 results on this basis is useful because it allows investors to more easily compare operating results for that year in a presentation comparable to the subsequent years presented in this report.

Table of Contents**Year ended December 31, 2003 compared to year ended December 31, 2002****Highlights**

The earnings increase in 2003 is characterized by a stabilization of the operating margins. This was a result of two developments:

improving operating margin in North America. After significant investments into our UltraCare program, which included the conversion to single-use dialyzers, the program now provides returns which contributed to an improvement of the operating margin in North America from 13.1% in 2002 to 13.8% in 2003.

price pressure in Germany, impact from the politically unstable situation in the Middle East and changes in the distribution system in Asia Pacific which led to a reduction of the operating margins in the International segment from 16.6% in 2002 to 15.2% in 2003.

During 2003, we reached settlements on all litigation relating to activities involving W.R. Grace before the 1996 Merger. We believe that the 2001 special charge for legal matters is sufficient to cover all related costs.

Cash flow provided from operations reached \$754 million and exceeded the prior year's cash flow from operations by \$204 million. This favorable development is a result of our focus on receivable collections and \$132 million of temporary liquidity provided by hedging of certain inter-company financing transactions, which is not expected to reoccur in that magnitude in 2004.

Consolidated Financials**Key Indicators for Consolidated Financials**

	2003	2002	Change in %	
			as reported	at constant exchange rates
Number of treatments	17,821,185	16,383,615	9%	
Same store treatment growth in %	4.9%	4.8%		
Revenue in \$ million	5,528	5,084	9%	5%
Gross profit in % of revenue	33.1%	32.6%		
Selling, general and administrative costs in % of revenue	18.5%	18.0%		
Net income in \$ million	331	290	14%	10%

Net revenue increased for the year ended December 31, 2003 over the comparable period in 2002 due to growth in revenue in both dialysis care and dialysis products.

Dialysis care revenue grew by 7% to \$3,978 million (6% at constant exchange rates) in 2003 mainly due to the growth in same store treatments, combined with acquisitions and the transition of billing for Medicare peritoneal dialysis patients from Method II billing to Method I billing. In 2002, peritoneal dialysis patients in the United States were billed by our products division (Method II) for their treatments. Beginning on January 1, 2003, they were billed by our services division (Method I). Dialysis product revenue increased by 13% to \$1,549 million (3% at constant exchange rates) in the same period.

Gross profit margin improved to 33.1% in the year ended December 31, 2003 from 32.6% for 2002. The increase is primarily a result of reduced dialysis care operating costs and dialysis product margin improvements in North America partially offset by the lower margin in the International segment. Depreciation and amortization expense for 2003 was \$217 million compared to \$211 million in 2002.

Approximately 43% of the Company's worldwide revenues are paid by and subject to regulations under governmental health care programs, primarily Medicare and Medicaid, administered by the United States government in both 2003 and 2002, respectively.

Selling, general and administrative costs increased from \$914 million in 2002 to \$1,022 million in 2003. Selling, general and administrative costs as a percentage of sales increased from 18.0% in 2002 compared to

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18.5% in 2003. This was in part due to the one time pension curtailment gain of \$12.6 million which reduced our selling, general and administrative costs in 2002. The remaining increase is mainly due to growth in international regions which have higher selling, general and administrative expenses partially offset by \$19 million of amortization expense for certain patient relationships and other intangible assets acquired in the 1996 Merger which were fully amortized in the fourth quarter of 2002. Net income for the period was \$331 million compared to \$290 million in 2002. Net income in 2002 was impacted by the \$12 million loss attributable to the early redemption of trust preferred securities.

In 2003, 17.8 million treatments were provided. This represents an increase of 9% over the same period in 2002. Same store treatment growth was 5% with additional growth of 3% from acquisitions. The remaining 1% increase in dialysis treatments was due to the transition of peritoneal dialysis patients from Method II (dialysis products) to Method I (dialysis service) billing in North America.

At December 31, 2003 we owned, operated or managed 1,560 clinics compared to 1,480 clinics at the end of 2002. During 2003, we acquired 42 clinics, opened 76 clinics and combined 38 clinics. The number of patients treated in clinics that we own, operate or manage increased from approximately 112,200 at December 31, 2002 to 119,250 at December 31, 2003. Average revenue per treatment for world-wide dialysis services decreased from \$226 to \$223 mainly due to the transition of peritoneal patients from Method II billing (dialysis products) to Method I (dialysis services).

The following discussions pertain to our business segments and the measures we use to manage these segments.

North America Segment**Key Indicators for North America Segment**

	<u>2003</u>	<u>2002</u>	<u>Change in %</u>
Number of treatments	12,366,028	11,638,740	6%
Same store treatment growth in %	3.8%	3.6%	
Revenue in \$ million	3,855	3,748	3%
EBITDA in \$ million	652	630	3%
EBITDA margin in %	16.9%	16.8%	
Depreciation and amortization in \$ million	120	139	-14%
Operating income in \$ million	532	491	8%
Operating income margin in %	13.8%	13.1%	

Revenue

Net revenue for the North America segment for the year ended December 31, 2003 grew in 2003 because dialysis care revenue increased by 4% from \$3,293 to \$3,429 million. This was partially offset by a decrease in product sales.

The increase in dialysis care revenue was driven by a 6% increase in treatments. Same store treatment growth was 4% and 1% resulted from acquisitions. A further 2% increase in dialysis treatments was due to a transition of peritoneal dialysis patients from Method II (dialysis products) to Method I (dialysis services). This was offset by a 1% decrease in treatments lost from clinics that were sold or closed and one less treatment day in 2003 compared to 2002. For this year the administration of EPO represented approximately 23% of total revenue.

At the end of 2003, approximately 82,400 patients were being treated in the 1,110 clinics that we own, operate or manage in the North America segment, compared to approximately 79,600 patients treated in 1,080 clinics at the end of 2002. The average revenue per treatment excluding laboratory testing revenue decreased from \$274 in 2002 to \$267 in 2003. Including laboratory testing the average revenue per treatment decreased from \$285 in 2002 to \$278 during 2003. This was mainly due to the transfer of our Method II patients to Method I.

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Dialysis product sales in both 2003 and 2002 include the sales of machines to a third-party leasing company which are leased back by our dialysis services division. Dialysis product sales in 2002 also includes Method II peritoneal dialysis revenues for our dialysis services patients. Method II patients were transferred to Method I effective January 1, 2003. Therefore there were no similar Method II revenues recorded in 2003. This reclassification of patients was the main cause of a 6% decrease in dialysis product revenue from \$454 million in 2002 to \$426 million in 2003. This was offset by an increase of sales due to the acquisition of the adsorber business of Fresenius AG in 2003. Our dialysis products division measures its external sales performance based on its sales to the net available external market. The net available external market excludes machine sales to third parties for machines utilized in the services division and Method II revenues involving our dialysis services division as well as sales to other vertically integrated dialysis companies and sales related to the adsorber business. Net available external market sales increased by 4% in 2003 over the comparable period 2002. The detail is as follows:

	Year ended December 31, 2003	Year ended December 31, 2002
	(in millions)	
Dialysis product sales	426	454
less sales to other vertically integrated dialysis companies and to leasing company of dialysis machines leased back	(34)	(42)
less method II and other		(37)
less sales related to adsorber business	(3)	
	—	—
Product sales to available external market	389	375

EBITDA.

EBITDA margin increased by 0.1%. This improvement in the margin is mainly a result of completion of the single-use dialyzer conversion which resulted in a reduction of dialysis care operating costs and an increase in product margin. Previous periods had been adversely affected by implementation costs of the single-use dialyzer program. This was partially offset by the pension curtailment gain of \$12.6 million in 2002.

Operating income

The increase in the operating margin was caused by lower depreciation and amortization as a result of the completion of amortization relating to patient relationships and other intangible assets acquired in the 1996 merger with an estimated useful life ending in the fourth quarter of 2002 and by the same factors causing the increase in the EBITDA margin stated above.

International Segment**Key Indicators for International Segment**

	2003	2002	Change in %	
			as reported	at constant exchange rates
Number of treatments	5,455,157	4,744,875	15%	
Same store treatment growth in %	7.7%	8.1%		
Revenue in \$ million	1,673	1,336	25%	11%
EBITDA in \$ million	349	292	20%	2%
EBITDA margin in %	20.8%	21.8%		
Depreciation and amortization in \$ million	95	70	37%	18%
Operating income in \$ million	254	222	14%	-4%
Operating income margin in %	15.2%	16.6%		

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Revenue

The increase in net revenues for the International segment resulted from increases in both dialysis care and dialysis product revenues. Acquisitions contributed approximately \$53 million (4%). Organic growth during the period was 7% (\$90 million) at constant exchange rates. This increase was improved by a \$193 million (14%) exchange rate effect due to the continued strengthening of the euro against the dollar in 2003.

Total dialysis care revenue increased by 32% (18% at constant exchange rates) to \$550 million in 2003 from \$416 million the same period of 2002. This increase is a result of base business growth of \$40 million combined with \$36 million in growth from acquisitions improved by approximately \$58 million due to exchange rate fluctuations.

As of December 31, 2003, approximately 36,850 patients were being treated at 450 clinics that we own, operate or manage in the International segment compared to 32,600 patients treated at 400 clinics at December 31, 2002. The average revenue per treatment increased from \$88 to \$101 (\$90 at constant exchange rates) due to the strengthening of the local currencies against the U.S. dollar and increased reimbursement rates partially offset by growth in countries with reimbursement rates below the average.

Total dialysis product revenue for 2003 increased by 22% (7% at constant exchange rates) to \$1,123 million. Including the effects of the acquisitions, the European region revenue increased \$272 million, a 30% increase (10% increase at constant exchange rates), the Latin America region revenue increased \$36 million or 24% (30% at constant exchange rates), while the Asia Pacific region revenue increased \$28 million or 10% (4% at constant exchange rates).

EBITDA.

Our EBITDA margin decreased from 21.8% to 20.8%. The main causes of this were price pressure in Europe, especially related to reimbursement changes in Germany which came into effect in the middle of 2003, increased cost of revenue due to the strengthening of the euro, lost revenues due to political instability in the Middle East and changes in the distribution system in Asia Pacific offset by retroactive reimbursement rate increases in Italy, Portugal and Venezuela.

Operating income

Our operating income margin decreased from 16.6% to 15.2%, due to the factors responsible for the decrease of EBITDA margin described above and higher depreciation and amortization mainly as a result of the expansion of production facilities in Europe and Asia Pacific.

Latin America

Our subsidiaries in Latin America contributed approximately 3% of our worldwide revenue and approximately 1% of our operating income in 2003. Our operations in Latin America were affected by the financial crisis and currency devaluations in nearly all currencies in Latin America whereas the Argentine Peso has recovered slightly. Because of these issues, we are experiencing lower than anticipated reimbursement rates, margin pressure and foreign currency exchange losses. In addition, the start-up of production and the entry into the peritoneal dialysis market in Mexico had an adverse effect on our margin in 2003.

In 2003, sales in Latin America increased 24% (30% at constant exchange rates) and operating income increased 21% (17% at constant exchange rates) compared to 2002. A worsening of the crisis in Latin America, a further devaluation of the Latin American currencies against the U.S. dollar or other unfavorable economic developments in Latin America, could result in an impairment of long lived assets and goodwill.

Corporate

We do not allocate corporate costs to our segments in calculating segment operating income and EBITDA as we believe that these costs are not within the control of the individual segments. These corporate costs

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primarily relate to certain headquarters overhead charges including accounting and finance, professional services, etc.

Total corporate operating loss was \$(29) million in the year ended December 31, 2003 compared to \$(18) million in the same period of 2002 to a large extent due to currency effects.

The following discussions pertain to our total Company costs.

Interest

Interest expense for 2003 decreased 6% compared to the same period in 2002 due to the charge recorded in the first quarter of 2002 for the redemption of trust preferred securities. See Note 13 *Mandatorily Redeemable Trust Preferred Securities* in our Consolidated Financial Statements.

Income Taxes

The effective tax rate for the year ended December 31, 2003 was 39.0% compared to 37.4% during the same period in 2002. This increase was caused by an increase of additional tax provisions and an increase in German tax rates in 2003.

Year ended December 31, 2002 compared year ended December 31, 2001

Net revenues for the year ended December 31, 2002 increased by 5% (6% at constant exchange rates) to \$5,084 million from \$4,859 million for the comparable period in 2001. The gross profit margin decreased from 33.7% to 32.6% in the year ended December 31, 2002 compared to the same period in 2001. This was mainly due to lower margins in North America and currency losses in Latin America (see also the discussion of our operating segments). Depreciation and amortization expense for 2002 was \$211 million compared to \$324 million for 2001. Amortization expense for goodwill and intangible assets not amortized anymore under SFAS No. 142 was \$121 million in 2001. Selling, general and administrative costs decreased from \$966 million in 2001 to \$914 million in 2002 due to the net of lower amortization mainly as a result of the adoption of SFAS No. 142 and higher bad debt expense and other operating expenses. Net income for the year was \$290 million as compared to \$63 million in 2001. The results of operations for 2002 reflect the implementation of SFAS No. 142 as of January 1, 2002, and for 2001 the special charge for legal matters. Income before extraordinary loss in 2002 was \$302 million compared to \$63 million in 2001. Earnings per Ordinary share before extraordinary loss for 2002 were \$3.12 compared to \$0.65 in 2001.

At December 31, 2002 we owned, operated or managed 1,480 clinics compared to 1,400 clinics at the end of 2001. During 2002, we acquired 33 clinics treating a total of 2,223 patients, opened 90 clinics and combined 43 clinics. The number of patients treated in clinics that we own, operate or manage increased from approximately 105,830 at December 31, 2001 to 112,200 at December 31, 2002. Approximately 16,385,000 treatments were provided in 2002; an increase of 7% from 15,250,000 treatments in 2001. Average revenue per treatment for world-wide dialysis services decreased from \$233 to \$226 mainly due to the decline of certain currencies compared to the U.S. dollar.

The following discussions pertain to our business segments and the measures we use to manage these segments.

North America Segment

Revenue

Net revenue for the North America segment for the year ended December 31, 2002 grew by 4% from \$3,602 million to \$3,748 million. Dialysis care revenue increased 5% from \$3,131 to \$3,293 million. The majority of the increase in dialysis care revenue resulted from a 4% increase in the number of treatments, mostly from same store growth, combined with a 1% increase due to increased revenue per treatment.

During 2002, approximately 79,600 patients were being treated in the 1,080 clinics that we own, operate or manage in the North America segment, compared to approximately 76,600 patients treated in 1,030 clinics during

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2001. The average revenue per treatment excluding laboratory testing revenue increased from \$273 in 2001 to \$274 in 2002. Including laboratory testing the average revenue per treatment increased from \$284 in 2001 to \$285 during 2002. The Medicare reimbursement rate has not increased since April 1, 2001, when it was increased by 1.6%. Medicare and Medicaid account for over 66% of North America dialysis services revenue.

Dialysis products revenue decreased 4% from \$471 million to \$454 million. Our North America dialysis products division measures its sales performance based on its sales to the net available external market which is in essence patients and non-vertically integrated third party dialysis providers. The net available external market excludes certain product revenues relating to our dialysis services business and sales to other vertically integrated dialysis companies. Comparing 2002 to 2001, sales to the net available external market increased 6%. The details are as follows:

	Year ended December 31, 2002	Year ended December 31, 2001
	(in millions)	
Dialysis product sales	454	471
less sales to other vertically integrated dialysis companies and to leasing company of dialysis machines leased back	(42)	(72)
less method II and other	(37)	(46)
	<u>375</u>	<u>353</u>
Product sales to available external market	375	353

EBITDA.

EBITDA for the North America segment decreased by 9% from \$693 million to \$630 million. The EBITDA margin decreased from 19.2% to 16.8%. The main reasons were expenses related to the implementation of our UltraCare program, an Amgen price increase for EPO, higher facilities lease and certification expenses and higher bad debt expense. In 2002 we continued the implementation of UltraCare which uses, among others, the latest technology of single use, high-flux polysulfone dialyzers to improve patient outcomes and care. The shift to single use of dialyzers in UltraCare created a higher cost per treatment compared to re-use. See Item 4B- Information on the Company-Dialysis Care-Fresenius UltraCare Program. This extra cost was mitigated through lower personnel costs combined with cost savings on medical supplies, making the single use of dialyzers cost neutral once implementation was completed. The implementation of this program had a negative effect on our margin since the additional expenses were incurred immediately, whereas the expected cost savings have been achieved over a longer period. A one-time pension curtailment gain was partially offset by severance and payroll costs for workforce reductions.

Depreciation and Amortization.

Depreciation and amortization decreased from 7% (\$247 million) of revenue in 2001 to 4% (\$139 million) in 2002. The decrease in amortization expense of \$108 million related almost exclusively to the accounting change required under SFAS No. 142. Adjusting the year ended December 31, 2001 as if SFAS No. 142 was implemented on January 1, 2001, amortization and depreciation remained at about 4% of revenue for both 2001 and 2002.

Operating Income

Operating income for the North America segment increased by 10%, from \$446 million to \$491 million due to the elimination of amortization for goodwill and intangible assets with indefinite useful lives under SFAS No. 142, partially offset by the factors affecting EBITDA. Assuming SFAS No. 142 had been adopted as of January 1, 2001, operating income would have decreased 11%, from \$553 million to \$491 million. The operating income margin over the same period decreased from 15.4% to 13.1% for the same reasons as described in EBITDA.

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International Segment

Revenue

Net revenue for the International segment during 2002 grew by 6% (12% at constant exchange rates) from \$1,257 million in 2001 to \$1,336 million in 2002. Acquisitions contributed approximately \$51 million (4%). Same store growth during the period was 8% (\$104 million). These increases in revenue were offset by a \$75 million (6%) adverse exchange rate effect, principally attributable to currency devaluation in Argentina. Including the effects of acquisitions, Asia Pacific region revenue increased \$53 million or 25% (26% at constant exchange rates), Latin America region revenue decreased \$95 million or 39% (an 11% increase at constant exchange rates) while European region revenue increased \$121 million, a 15% increase (9% increase at constant exchange rates).

Total dialysis care revenue decreased during 2002 by 2% (a 19% increase at constant exchange rates) to \$416 million from \$426 million the same period of 2001. This decrease is a result of base business growth of \$42 million combined with \$40 million in growth from acquisitions offset by approximately \$92 million due to exchange rate fluctuations.

As of December 31, 2002, approximately 32,600 patients were being treated at 400 clinics that we own, operate or manage in the International segment compared to 29,230 patients treated at 370 clinics at December 31, 2001. The average revenue per treatment decreased from \$104 to \$88 due to the depression of local currencies against the U.S. dollar. At constant exchange rates, revenue per treatment increased \$3 to \$107.

Total dialysis product revenue for 2002 increased by 11% (9% at constant exchange rates) to \$921 million. Sales of dialyzers and peritoneal dialysis equipment more than offset a decrease in hemodialysis machine sales.

EBITDA.

EBITDA for the International segment was \$292 million for both 2002 and 2001 (a decrease of 1% at constant exchange rates). Our EBITDA margin decreased from 23.2% to 21.8% mainly due to the financial crisis in Latin America combined with growth of our business in lower margin countries.

Depreciation and Amortization

Depreciation and amortization decreased slightly from 6% (\$76 million) to 5% (\$70 million) of revenues for 2002 compared to 2001 mainly as a result of the implementation of SFAS No. 142, partially offset by additional depreciation and amortization for expanded production facilities in Europe and Asia Pacific. 2001 amortization expense for goodwill and intangible assets not amortized in 2002 under SFAS No. 142 amounted to \$14 million.

Operating Income

Operating income for the International segment for 2002 increased 3% (a 1% decrease at constant exchange rates) to \$222 million due to the implementation of SFAS No. 142. Our operating income margin decreased slightly from 17.2% to 16.6%. Adjusting 2001 as if SFAS No. 142 was implemented on January 1, 2001, operating income decreased 3% (a decrease of 6% at constant exchange rates) with operating income margin decreasing from 18.3% to 16.6%. As with EBITDA, this decrease was caused mainly by the financial crisis in Latin America and growth in lower margin countries.

Latin America

Our subsidiaries in Latin America contributed approximately \$149 million (3%) of our worldwide revenue in 2002 compared to approximately \$244 million (5%) of our worldwide revenue in 2001. EBITDA decreased from \$36 million in 2001 to \$14 million in 2002 while operating income decreased from \$21 million to \$6 million in the same period. Our operations in Latin America were affected by the financial crisis and the currency devaluation in Argentina and other Latin America countries.

We considered the financial crisis in Latin America a triggering event. In the third quarter of this year, we completed an impairment test of our Latin America operations as required by SFAS No. 142. As of

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September 30, 2002, there was no impairment of long lived assets and goodwill. However, a worsening of the crisis in Argentina, a further devaluation of the Argentine peso or other Latin American currencies against the U.S. dollar or other unfavorable economic developments in Latin America could result in an impairment of long lived assets and goodwill. Goodwill and long-lived assets amount to \$26 million and \$84 million, respectively, at December 31, 2002.

Corporate

We do not allocate corporate costs to our segments in calculating segment operating income and EBITDA as we believe that these costs are not within the control of the individual segments. These corporate costs primarily relate to certain headquarters overhead charges including accounting and finance, professional services, etc.

Total corporate operating income was \$(18) million in 2002 compared to \$(25) million in 2001. Operating income improved in 2002 due to a one-time recognition of \$7 million in 2001 expenses related to 1996 Merger related legal matters.

On February 14, 2002, the Company redeemed the entire \$360 million aggregate amount outstanding of its 9% Trust Preferred Securities due 2006. The Company exercised its option to redeem the securities at a price of US\$1,045 per \$1,000 liquidation amount plus accrued distributions of \$18.25 per \$1,000 for a total redemption price of \$1,063.25 per \$1,000. The Company funded the redemption utilizing its 1996 Senior Credit Agreement.

An extraordinary loss of \$12 million was incurred as a result of the early redemption of debt, consisting of \$16 million of redemption premiums plus \$3 million of associated debt issuance costs, less a \$7 million tax benefit.

The following discussions pertain to our total Company costs.

Interest

Net interest expense for 2002 decreased 7% compared to 2001 mainly due to the redemption of our 9% trust preferred securities in February 2002, which we financed through our credit agreement at lower rates.

Income Taxes

The effective tax rate for the year ending December 31, 2002 was 37.5% compared to 58.4% during 2001. This decrease in the effective rate was caused by the elimination of non-deductible amortization expense due to SFAS No. 142 combined with not being able to deduct a portion of the special charge for legal matters in 2001.

B. LIQUIDITY AND CAPITAL RESOURCES

Year ended December 31, 2003 compared to year ended December 31, 2002

Cash Flow

Operations

We generated cash from operating activities of \$754 million in 2003 and \$550 million in the comparable period in 2002, an increase of approximately 37% over the prior year. Cash flows benefited from \$132 million of temporary liquidity provided by hedging of certain intercompany financing transactions, which is not expected to reoccur in that magnitude in 2004, improved accounts receivable collections and lower prepaid expenses and other current assets. We classify the cash outflows from our accounts receivable securitization program in the amount of \$287 million as a financing activity.

Investing

Cash used in investing activities increased from \$281 million to \$369 million mainly because of increased purchases of property, plant and equipment. Capital expenditures for property, plant and equipment net of disposals were \$276 million for the year ended December 31, 2003 and \$201 million for the comparable period in

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2002. In 2003, capital expenditures were \$170 million in the North America segment and \$106 million for the International segment. In 2002, capital expenditures were \$98 million in the North America segment and \$103 million for the International segment. The majority of our capital expenditures were used for equipment in new clinics, the buyout of the Ogden lease, improvements to existing clinics, and expansion of production facilities. Net capital expenditures were approximately 5% of total revenue.

In 2003, we paid approximately \$92 million (\$40 million for the North American segment and \$52 million for the International segment) cash for acquisitions consisting primarily of the adsorber business acquired from Fresenius AG and dialysis clinics. In accordance with the requirements of the pooling agreements relating to outstanding Ordinary shares and Preference shares, the acquisition of the Fresenius AG adsorber business was approved by our independent directors. See Item 10, *Additional Information* Description of the Pooling Agreements. In the same period in 2002, we paid approximately \$80 million (\$38 million for the North American segment and \$42 million for the International segment) cash for acquisitions consisting primarily of dialysis clinics.

Financing

Net cash used in financing was \$416 million in 2003 compared to \$265 million in the same period of 2002. Our financing needs decreased due to higher operating cash flow partially offset by higher payments for investing activities, higher dividend payments and payments for the redemption of the FMCH Class D Preferred Stock. Cash on hand was \$48 million at December 31, 2003 compared to \$65 million at December 31, 2002.

On February 21, 2003, we entered into an amended and restated bank agreement with Bank of America N.A, Credit Suisse First Boston, Dresdner Bank AG New York, JPMorgan Chase Bank, The Bank of Nova Scotia and certain other lenders (collectively, the *Lenders*), pursuant to which the Lenders have made available to the Company and certain subsidiaries and affiliates an aggregate amount of up to \$1.5 billion through three credit facilities. On August 22, 2003, the 2003 Senior Credit Agreement was amended so that, in effect, the aggregate amount of \$1.5 billion was voluntarily reduced to \$1.4 billion and the interest rate on a new term loan facility (Loan C) was 25 basis points lower than the interest rate on Loan B which was repaid. Funds available under this agreement were used to refinance the previous credit agreement's outstanding balances and to pay down \$287 million of our accounts receivable facility.

On March 28, 2003, FMCH redeemed all of its outstanding shares of Class D Special Dividend Preferred Stock (*Class D Shares*) at a total cash outflow of approximately \$9 million.

On February 14, 2002, we redeemed the entire \$360 million amount outstanding of our 9% Trust Preferred Securities due 2006, utilizing funds borrowed under our 1996 senior credit agreement. A loss of \$12 million after tax was incurred as a result of the early redemption of debt, consisting of \$16 million of redemption premiums plus a \$4 million write-off of associated debt issuance costs, less a \$8 million tax benefit.

Further financing was provided by Fresenius AG at different levels throughout the year. As of December 31, 2003 the balance outstanding was \$30 million.

Dividends

Consistent with prior years, we will continue to follow an earnings-driven dividend policy. The Managing Board and the supervisory board will propose a dividend of 1.02 per ordinary share (2002: 0.94) and 1.08 per preference share (2002: 1.00) for shareholder approval at the annual general meeting on May 27, 2004. The total expected dividend payment is approximately 99.7 million. Our 2003 Senior Credit Agreement limits disbursement of dividends and other restricted payments during 2004 to \$150 million.

Liquidity

Our primary sources of liquidity have historically been cash from operations, cash from short-term borrowings as well as from long-term debt from third parties and from related parties and cash from issuance of Preference shares and trust preferred securities. Cash from operations is impacted by the profitability of our business and the development of our working capital, principally receivables. The profitability of our business

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depends significantly on reimbursement rates. Approximately 72% of our revenues are generated from providing dialysis treatment, a major portion of which is reimbursed by either public health care organizations or private insurers. For the year ended December 31, 2003, approximately 43% of our consolidated revenues resulted from U.S. federal health care benefit programs, such as Medicare and Medicaid reimbursement. Legislative changes may affect all Medicare reimbursement rates for the services we provide, as well as the scope of Medicare coverage. A decrease in reimbursement rates could have a material adverse effect on our business, financial condition and results of operations and thus on our capacity to generate cash flow. Furthermore cash from operations depends on the collection of accounts receivable. We may face difficulties in enforcing and collecting accounts receivable under some countries' legal systems. Some customers and governments may have longer payment cycles. This could have a material adverse effect on our capacity to generate cash flow.

Cash from short-term borrowings can be generated by selling interests in accounts receivable (accounts receivable facility) and by borrowing from our parent Fresenius AG. Long-term financing is provided by the revolving portion and term loans under our 2003 Senior Credit Agreement and has been provided through the issuance of our trust preferred securities. We believe that our existing credit facilities, cash generated from operations and other current sources of financing are sufficient to meet our foreseeable needs.

The proceeds of Loan C, together with cash from operations, were used to voluntarily and permanently repay Loan B, a \$500 million term loan facility under the 2003 Senior Credit Agreement. We used the initial borrowings under the 2003 Senior Credit Agreement to refinance outstanding borrowings under our prior senior credit agreement and to repay \$287 million of the accounts receivable facility.

At December 31, 2003, we had approximately \$463 million of borrowing capacity available under the revolving portion of our 2003 Senior Credit Agreement.

Our Senior Credit Agreement and the indentures relating to our trust preferred securities include covenants that require us to maintain certain financial ratios or meet other financial tests. Under our 2003 Senior Credit Agreement, we are obligated to maintain a minimum consolidated net worth and a minimum consolidated fixed charge ratio (ratio of earnings before interest, taxes, depreciation, amortization and rent to fixed charges) and we have to maintain a certain consolidated leverage ratio (ratio of consolidated funded debt to adjusted EBITDA).

Our 2003 Senior Credit Agreement and our indentures include other covenants which, among other things, restrict or have the effect of restricting our ability to dispose of assets, incur debt, pay dividends and other restricted payments (limited to \$130 million in 2003, increasing to \$150 million in 2004), create liens or make capital expenditures, investments or acquisitions. The breach of any of the covenants could result in a default under the 2003 Senior Credit Agreement or the notes, which could, in turn, create additional defaults under the agreements relating to our other long-term indebtedness. In default, the outstanding balance under the 2003 Senior Credit Agreement becomes due at the option of the Lenders. As of December 31, 2003, we are in full compliance with all financial covenants under the 2003 Senior Credit Agreement.

After redemption of \$360 million aggregate liquidation amount of 9% trust preferred securities on February 14, 2002, our long-term financing under our remaining trust preferred securities begins to come due in February 2008. However, Loan C under our amended 2003 Senior Credit Agreement will become due on October 31, 2007 if our trust preferred securities due February 1, 2008 are not repaid or refinanced or their maturity is not extended prior to that date.

National Medical Care, Inc. (NMC), our subsidiary, has an asset securitization facility (the accounts receivable facility) whereby receivables of NMC and certain affiliates are sold to NMC Funding Corporation (the Transferor), a wholly-owned subsidiary of NMC, and subsequently the Transferor transfers and assigns percentage ownership interests in the receivables to certain bank investors. The amount of the accounts receivable facility was last amended on October 23, 2003, when we extended its maturity to October 22, 2004. Funds from the 2003 Senior Credit Agreement were used to pay down \$287 million of the accounts receivable facility in 2003.

Our capacity to generate cash from the accounts receivable facility depends on the availability of sufficient accounts receivable that meet certain criteria defined in the agreement with the third party funding corporation. A

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lack of availability of such accounts receivable could have a material impact on our capacity to utilize the facility for our financial needs.

The settlement agreement with the asbestos creditors committees on behalf of the W.R. Grace & Co. bankruptcy estate provides for payment of \$115 million upon approval of the settlement agreement by the U.S. District Court, which has occurred, and confirmation of a W.R. Grace & Co. bankruptcy reorganization plan that includes the settlement.

We are subject to a tax audit in Germany and as a result may be required to make additional tax payments. The potential payments will not affect earnings, as the related taxes have been fully accrued. We are currently not in a position to determine the timing of these payments which may become payable in 2004.

Obligations

The following table summarizes, as of December 31, 2003, our obligations and commitments to make future payments under our long-term debt, trust preferred securities and other long term obligations, and our commitments and obligations under lines of credit and letters of credit.

Contractual Cash Obligations	Total	Payments due by period of		
		1 Year	2-5 Years	Over 5 Years
		(in millions)		
Trust Preferred Securities	\$ 1,242	\$ 644	\$ 598	
Long Term Debt	1,192	86	709	397
Capital Lease Obligations	10	4	5	1
Operating Leases	1,091	237	589	265
Unconditional Purchase Obligations	220	94	126	
Other Long-term Obligations	5	5		
	<u>\$ 3,760</u>	<u>\$ 426</u>	<u>\$ 2,073</u>	<u>\$ 1,261</u>

Available Sources of Liquidity	Total	Expiration per period of		
		1 Year	2-5 Years	Over 5 Years
Unused Senior Credit Lines	\$ 463	\$ 463		
Other Unused Lines of Credit	96	96		
	<u>\$ 559</u>	<u>\$ 96</u>	<u>\$ 463</u>	<u>\$</u>

The amount of guarantees and other commercial commitments at December 31, 2003 is not significant.

Borrowings

Short-term borrowings of \$89 million and \$125 million at December 31, 2003, and 2002, respectively, represent amounts borrowed by certain of our subsidiaries under lines of credit with commercial banks. The average interest rates on these borrowings at December 31, 2003, and 2002 was 3.38% and 4.67%, respectively. For information regarding short-term borrowings from affiliates see Note 4b in our Consolidated Financial Statements.

Excluding amounts available under the 2003 Senior Credit Agreement (as described below), at December 31, 2003, we had \$96 million available under such commercial bank agreements. Some of these lines of credit are secured by the individual borrowers' accounts receivable and contain various covenants including, but not limited to, requirements for maintaining defined levels of working capital, net worth, capital

expenditures and certain financial ratios.

On February 21, 2003, we entered into an amended and restated senior credit agreement with Bank of America N.A, Credit Suisse First Boston, Dresdner Bank AG New York, JPMorgan Chase Bank, The Bank of Nova Scotia, and certain other financial institutions (collectively, the Lenders). Pursuant to the agreement, the Lenders made available to the Company and certain subsidiaries and affiliates an aggregate amount of up to

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\$1.5 billion through three credit facilities. On August 22, 2003 the 2003 Senior Credit Agreement was amended so that, in effect, the aggregate amount was voluntarily reduced to \$1.4 billion and the interest rate on a new term loan facility (Loan C) was 25 basis points lower than Loan B, which was repaid. The credit facilities are a revolving facility of \$500 million a term loan facility of \$500 million (Loan A) and a term loan facility of \$400 million (Loan C).

In 2001, we issued four tranches of senior notes (Euro Notes) totaling 128.5 million. The first tranche was for 80 million with a fixed interest rate of 6.16% and the second and third tranches for 28 million and 15 million, respectively, with variable interest rates which averaged 3.84% in 2003 and 4.78% in 2002. The final tranche was for 5 million at a fixed rate of 5.33%. All four tranches have a maturity date of July 13, 2005. Both floating rates are tied to the EURIBOR rate.

Year ended December 31, 2002 compared to year ended December 31, 2001

Cash Flow

Operations

We generated cash from operating activities of \$550 million in the year ended December 31, 2002 and \$424 million in the comparable period in 2001, an increase of about 30% over the prior year. Cash flows benefited from improved accounts receivable collections, especially in North America.

Investing

Cash used in investing activities decreased from \$468 million to \$281 million mainly because of lower cash acquisition payments and net capital expenditures. In 2002, we paid approximately \$80 million (\$38 million for the North American segment and \$42 million for the International segment) cash for acquisitions consisting primarily of dialysis clinics. In 2001, we paid approximately \$217 million (\$178 million for the North American segment and \$39 million for the International segment) cash for acquisitions consisting primarily of dialysis clinics, including the cash portion of the purchase price for Everest.

In addition, capital expenditures for property, plant and equipment net of disposals were \$201 million for the year ended December 31, 2002 and \$251 million for the comparable period in 2001. In 2002, capital expenditures were \$98 million in the North America segment and \$103 million for the International segment. In 2001, capital expenditures were \$123 million in the North America segment and \$128 million for the International segment. The majority of our capital expenditures were used for the upgrading of existing clinics and the expansion of production facilities in North America, Germany, France, Italy, Mexico, and Brazil. Capital expenditures were approximately 4% of total revenue.

Financing

Net cash used in financing was \$265 million in 2002 compared to cash provided by financing of \$43 million in 2001 because our financing needs decreased due to lower borrowing for acquisitions, higher operating cash flows and lower capital expenditure. Cash on hand was \$65 million at December 31, 2002 compared to \$62 million at December 31, 2001.

On February 14, 2002, we redeemed the entire \$360 million aggregate liquidation amount outstanding of our 9% Trust Preferred Securities due 2006, utilizing funds borrowed under our 1996 senior credit facility. An extraordinary loss of \$12 million was incurred as a result of the early redemption of debt, consisting of \$16 million of redemption premiums plus a \$3 million write-off of associated debt issuance costs, less a \$7 million tax benefit.

In January 2001, we completed the acquisition of Everest. Approximately one-third of the purchase price (\$365 million) was funded by the issuance of 2.25 million Preference shares (\$99 million) to Everest stockholders. The remaining purchase price was paid with \$131 million cash and debt assumed (\$135 million). This debt was subsequently retired using our senior credit facility.

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In June 2001 we completed offerings of \$225 million aggregate liquidation amount of dollar-denominated 7 7/8% Trust Preferred Securities due 2011 and 300 million aggregate liquidation amount of euro-denominated 7 3/8% trust preferred securities due 2011.

Between July 13, 2001 and December 5, 2001 we issued four tranches of senior notes totaling 128.5 million.

Dividends

Consistent with prior years, in 2002 we followed an earnings driven dividend policy. The management board and supervisory board proposed a dividend of 0.94 per ordinary share (2001: 0.85) and 1.00 per preference share (2001: 0.91) to shareholders at the annual general meeting on May 22, 2003 and they were approved. The total dividend payment was approximately 92 million. Our Senior Credit Agreement limited dividend payments during 2003 to \$130 million.

Liquidity

Primary sources of liquidity have historically been cash from operations, cash from short term borrowings as well as from long term debt from third parties and from related parties and cash from issuance of Preference shares. We expect that our primary source of liquidity for 2003 will be our operations and financing activities. Cash from operations is impacted by the profitability of our business and the development of our working capital, principally receivables. The profitability of our business depends significantly on reimbursement rates. Approximately 73% of our revenues are generated from providing dialysis treatment, a major portion of which is reimbursed by either public health care organizations or private insurers. For the twelve months ended December 31, 2002, approximately 43% of our consolidated revenues resulted from U.S. federal health care benefit programs, such as Medicare and Medicaid reimbursement. Legislative changes may affect all Medicare reimbursement rates for the services we provide, as well as the scope of Medicare coverage. A decrease in reimbursement rates could have a material adverse effect on our business, financial condition and results of operations and thus on our capacity to generate cash flow. For additional information about the impact of U.S. government reimbursement programs, see Item 4B, Business Overview Dialysis Care Sources of U.S. Dialysis Care Net Revenues. Furthermore cash from operations depends on the collection of accounts receivable. We may face difficulties in enforcing and collecting accounts receivable under some countries' legal systems. Some customers and governments may have longer payment cycles. This could have a material adverse effect on our capacity to generate cash flow.

Cash from short-term borrowings can be generated by selling interests in accounts receivable (accounts receivable facility) and by borrowing from our parent Fresenius AG. Long-term financing is provided by the revolving portion and term loans of our senior credit facility and has been provided through the issuance of our trust preferred securities facility. We believe that our existing credit facilities, cash generated from operations and other current sources of financing are sufficient to meet our foreseeable needs.

At December 31, 2002, we had approximately \$381 million of borrowing capacity available under the revolving portion of our senior credit facility. On February 21, 2003, we entered into an amended and restated senior credit agreement with Bank of America N.A, Credit Suisse First Boston, Dresdner Bank AG New York, JPMorgan Chase Bank, The Bank of Nova Scotia, and certain other financial institutions. Pursuant to the agreement, the Lenders have made available to the Company and certain subsidiaries and affiliates an aggregate of up to \$1,500 million through three credit facilities. The three facilities are a revolving facility of \$500 million and two term loan facilities of \$500 million each. We used the initial borrowings under the 2003 Senior Credit Agreement to refinance outstanding borrowings under our prior senior credit agreement and for general corporate purposes (see note 25 of the Consolidated Financial Statements).

Our 2003 Senior Credit Agreement and the indentures relating to our trust preferred securities include covenants that require us to maintain certain financial ratios or meet other financial tests. Under our Senior Credit Agreement, we are obligated to maintain a minimum consolidated net worth and a minimum consolidated fixed charge ratio (ratio of earnings before interest, taxes, depreciation, amortization and rent to fixed charges) and we have to maintain a certain consolidated leverage ratio (ratio of consolidated funded debt to EBITDA).

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Our 2003 Senior Credit Agreement and our indentures include other covenants which, among other things, restrict or have the effect of restricting our ability to dispose of assets, incur debt, pay dividends, create liens or make capital expenditures, investments or acquisitions. The breach of any of the covenants could result in a default under the credit agreement or the notes, which could, in turn, create additional defaults under the agreements relating to our other long term indebtedness. In default, the outstanding balance under the senior credit agreement becomes due.

After redemption of \$360 million aggregate liquidation amount of 9% trust preferred securities on February 14, 2002, our long-term financing under our remaining trust preferred securities begins to come due in February 2008.

NMC, our subsidiary, has an asset securitization facility (the accounts receivable facility) whereby receivables of NMC and certain affiliates are sold to NMC Funding Corporation (the Transferor), a wholly-owned subsidiary of NMC, and subsequently the Transferor transfers and assigns percentage ownership interests in the receivables to certain bank investors.

Our capacity to generate cash from the accounts receivable facility depends on the availability of sufficient accounts receivable that meet certain criteria defined in the agreement with the third party funding corporation. A lack of availability of such accounts receivable may have a material impact on our capacity to utilize the facility for our financial needs.

The settlement agreement with the asbestos creditors committee on behalf of the W.R. Grace & Co. bankruptcy estate provides for payment of \$115 million upon confirmation of the W.R. Grace & Co. bankruptcy reorganization and approval of the settlement agreement by the U.S. Bankruptcy Court. We are subject to a tax audit in Germany and as a result may be required to make additional tax payments. The potential payments will not affect earnings, as the related taxes have been fully accrued. We are currently not in a position to determine the amount and timing of these payments.

Recently Issued Accounting Standards

In August 2001, the FASB issued SFAS No. 143, *Accounting for Asset Retirement Obligations*. SFAS No. 143 requires that the fair value of a liability for an asset retirement obligation be recognized in the period in which it is incurred if a reasonable estimate of fair value can be made. The associated asset retirement costs are capitalized as part of the carrying amount of the long-lived asset. It applies to legal obligations associated with the retirement of long-lived assets that result from the acquisition, construction, development and/or normal operation of a long-lived asset. We adopted SFAS No. 143 as of January 1, 2003. The adoption of SFAS No. 143 did not have a material impact on our financial statements.

In April 2002, the FASB issued SFAS No. 145, *Rescission of FASB Statements No. 4, 44, and 64, Amendment of FASB Statement No. 13, and Technical Corrections*. SFAS No. 145 rescinds SFAS No. 4, SFAS No. 64 related to classifications of gains and losses on debt extinguishments such that most debt extinguishment gains and losses will no longer be classified as extraordinary. SFAS No. 145 also amends SFAS No. 13, with respect to certain sale-leaseback transactions. We adopted SFAS No. 145 in regard to SFAS No. 4 on January 1, 2003. In the first quarter of 2002, we recorded an extraordinary loss of \$11.8 million, net of taxes of \$7.7 million, as a result of the early redemption of debt (see Note 13). This loss is no longer presented as an extraordinary loss upon the adoption of SFAS No. 145. We adopted the other provisions of SFAS No. 145 effective April 1, 2002.

In June 2002, the FASB issued SFAS No. 146, *Accounting for Costs Associated with Exit or Disposal Activities*. The standard requires companies to recognize costs associated with exit or disposal activities when liabilities are incurred. SFAS No. 146 replaces EITF Issue No. 94-3, *Liability Recognition for Certain Employee Termination Benefits and Other Costs to Exit an Activity (Including Certain Costs Incurred in a Restructuring)*. This statement is applied prospectively to exit or disposal activities initiated after December 31, 2002.

In November 2002, the Financial Accounting Standards Board issued FASB Interpretation No. 45 (FIN 45), *Guarantor's Accounting and Disclosure Requirements for Guarantees of Indebtedness of Others*. FIN 45 also requires the guarantor to recognize a liability for the non-contingent component of the guarantee, that is, the obligation to stand ready to perform in the event that special triggering events or conditions occur. The

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initial recognition and measurement provisions are applicable prospectively to guarantees issued or modified after December 31, 2002. FIN 45 also clarifies and expands the disclosure requirements related to guarantees, including product warranties. FIN 45 does not materially impact the Company's financial statements.

On April 3, 2003, the Financial Accounting Standards Board issued SFAS No. 149 *Amendment of Statement 133 on Derivative Instruments and Hedging Activities*. This Statement amends and clarifies financial accounting and reporting for derivative instruments, including certain derivative instruments embedded in other contracts (collectively referred to as derivatives) and for hedging activities under SFAS No. 133, *Accounting for Derivative Instruments and Hedging Activities*. This Statement is effective for contracts entered into or modified after June 30, 2003. This adoption did not have any impact on our financial statements.

In May 2003, the Financial Accounting Standards Board issued SFAS No. 150 *Accounting for certain Financial Instruments with Characteristics of both Liabilities and Equity*. This Statement requires an issuer to classify certain financial instruments with the characteristics of both liabilities and equity as a liability (or asset in some circumstances) instead of equity. This Statement is effective for financial instruments entered into or modified after May 31, 2003 and otherwise effective at the beginning of the first interim period beginning after June 15, 2003. This adoption did not have any impact on our financial statements.

In December 2003 the Financial Accounting Standards Board issued SFAS No. 132 (revised 2003) *Employers Disclosures about Pensions and Other Postretirement Benefits – an amendment of FASB Statements No. 87, 88 and 106*. This statement extends the publishing rules for pension liabilities according to SFAS No. 132. The accounting and valuation principles remain unchanged.

In December 2003, the Financial Accounting Standards Board issued FASB Interpretation No. 46R *Consolidation of Variable Interest Rate Entities (revised)* (FIN 46R) which addresses how a business enterprise should evaluate whether it has a controlling financial interest in an entity through means other than voting rights and accordingly should consolidate the entity. FIN 46R replaced FASB Interpretation No. 46 *Consolidation of Variable Interest Rate Entities* which was issued in January 2003. The Company is required to apply FIN 46R for special purpose entities as of December 31, 2003 and for all other Variable Interest Entities (VIEs) as of March 31, 2004. The Company is not involved with any special purpose entity which required initial consolidation as of December 31, 2003 and will apply FIN 46R on March 31, 2004 for all VIEs.

We are party to various arrangements with certain dialysis clinics to provide management services, financing and product supply. Some of these clinics are variable interest entities. Under FIN 46R these clinics will be consolidated if we are the primary beneficiary. We also participate in a joint venture which is engaged in the perfusion business. The arrangements with the joint venture partner are such that the joint venture qualifies as a variable interest entity and we are the primary beneficiary. These variable interest entities generate approximately \$153,000 in annual revenue. This includes approximately \$14,000 related to variable interest entities in which we are not the primary beneficiary. We have investments, other long term assets and receivables of approximately \$42,000 which represent our maximum exposure to loss as a result of our involvement with the variable interest entities.

C. Research and development

Our research and development activities aim to improve the quality of dialysis treatment by matching it more closely with the individual needs of the patient, while reducing the overall cost for treatment. With our vertical integration, our research and development department can apply our experience as the world's largest provider of dialysis treatments to product development. To maintain and further enhance a continuous stream of product innovations, we have 388 employees working in research and development worldwide at December 31, 2003. Approximately two-thirds of our research and development activities are based in Germany and one third in North America.

Research and development focuses strongly on the development of new products, technologies and treatment concepts to optimize treatment quality for dialysis patients, and on process technology for manufacturing our products. Research and development expenditures were \$36 million in 2001, \$47 million in 2002, and

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\$50 million in 2003. For information regarding recent product introductions, see Item 4.B. Business Overview New Product Introductions.

We intend to continue to maintain our central research and development operations for disposable products at our St. Wendel, Germany facility and for durable products at our Schweinfurt and Bad Homburg, Germany facilities. Local activities will continue to focus on cooperative efforts with those facilities to develop new products and product modifications for local markets.

In North America, we have concentrated our business development activities on expanding our products business in three main areas:

pharmaceutical products utilized in treating our renal patient base

innovative products to improve vascular access outcomes for our renal patients

products and technologies which leverage our core competencies to provide extracorporeal therapies to treat other diseases

D. Trend information

For information regarding significant trends in our business see Item 5.A. Operating Financial Review and Prospects.

Item 6. Directors, Senior Management and Employees

A. Directors and senior management

General

In accordance with the German Stock Corporation Act, we have a supervisory board and a management board. The two boards are separate and no individual may simultaneously be a member of both boards.

Our Supervisory Board

Our supervisory board consists of five members who are elected by the holders of Ordinary shares at our annual general meeting. Pursuant to the pooling agreements described in the exhibits, at least one-third (but no fewer than two) of the members of the supervisory board elected by the shareholders are required to be independent directors as defined in the pooling agreements, i.e., persons with no substantial business or professional relationship with us, Fresenius AG or any affiliate of either.

If and when either:

Fresenius Medical Care AG itself has more than 500 employees; or

we enter into a domination agreement with a German subsidiary having more than 500 employees, or if that subsidiary is integrated into Fresenius Medical Care AG;

the German employees of Fresenius Medical Care AG and our German subsidiaries will elect one-third of the members of the supervisory board. If and when the aggregate number of employees of Fresenius Medical Care AG and our German subsidiaries exceeds 2,000, the supervisory board will increase to 12 persons and the holders of Ordinary shares and the German employees of Fresenius Medical Care and our German subsidiaries will elect six members each. In that case, the Chairman of the supervisory board will be selected from the members elected by the shareholders and will have the tie-breaking vote.

The term of a member of the supervisory board will expire at the end of the general meeting of shareholders after the fourth fiscal year following the year in which the member was elected, but not counting the fiscal year in which such member's term begins. Members of the supervisory board elected by our shareholders may be removed by a resolution of our general meeting. This resolution requires a three-fourths majority of the votes cast at that meeting. The supervisory board ordinarily acts by simple majority vote and the Chairman has a tie-breaking vote in case of any deadlock.

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The principal function of the supervisory board is to appoint and to supervise the management board and to approve mid-term planning, dividend payments and matters which are not in the ordinary course of business and are of fundamental importance to us.

The table below provides the names and ages of the members of our supervisory board as of December 31, 2003.

Name	Age as of December 31, 2003
Dr. Gerd Krick, Chairman	65
Dr. Dieter Schenk, Deputy Chairman	51
Prof. Dr. Bernd Fahrholz	56
Walter L. Weisman ⁽¹⁾	68
Stephen M. Peck ⁽¹⁾	68

(1) Independent Director for purposes of our pooling agreement

The term for each member of the supervisory board set forth above will expire at the end of the fiscal year 2005. Members of the supervisory board are eligible for reelection at the 2006 Annual General Shareholders Meeting.

DR. GERD KRICK has been Chairman of our supervisory board since January 1, 1998. From 1992 to 2003, he was Chairman of the Fresenius AG management board. Prior to 1992, he was a Director of the Medical Systems Division of Fresenius AG and Deputy Chairman of the Fresenius AG management board. From September 1996 until December 1997, Dr. Krick was Chairman of the management board of Fresenius Medical Care. Dr. Krick is a member of the Board of Directors of Adelphi Capital Europe Fund, of the Administrative Board of Dresdner Bank Luxembourg S.A., of the supervisory board of Vereinte Krankenversicherung AG, of the Advisory Board of HDI Haftpflichtverband der deutschen Industrie and of the Board of Trustees of the Donau Universität Krems. He is also the Chairman of the supervisory boards of Vamed AG, Fresenius Kabi AG and Fresenius Kabi Austria GMBH all of which are subsidiaries of Fresenius AG.

DR. DIETER SCHENK has been Vice Chairman of our supervisory board since 1996. He is an attorney and tax advisor and has been a partner in the law firm of Nörr Stiefenhofer Lutz since 1986. Dr. Schenk is also a member of the supervisory board of Fresenius AG. He also serves as a member and chairman of the supervisory board of Gabor Shoes AG, a member and vice-chairman of the supervisory boards of Greiffenberger AG and TOPTICA Photonics AG.

PROF. DR. BERND FAHRHOLZ has been a member of our supervisory board since 1998. He is an attorney and was a member of the management board of Dresdner Bank AG since 1998 and its Chairman from April 2000 until he resigned in March of 2003. He also served as the deputy chairman of the management board of Allianz AG and chairman of the supervisory board of Advance Holding AG until March of 2003. Dr. Fahrholz was a member of the supervisory boards of Dresdner Kleinwort Benson North America Inc. until February of 2003, BNP-Paribas S.A. until March of 2003 and Dresdner Bank Luxembourg S.A. until October of 2003. He currently serves on the supervisory boards of BMW AG and Heidelberg Cement AG.

WALTER L. WEISMAN has been a member of our supervisory board since 1996. He is a private investor and a former Chairman and Chief Executive Officer of American Medical International, Inc. Mr. Weisman is on the board of Community Care Health Network, Inc., Maguire Properties, Inc., and Occidental Petroleum Corporation. He is Vice-Chairman of the Board of trustees for the California Institute of Technology, Chairman of the Board of the Los Angeles County Museum of Art, Chairman of the Board of the Sundance Institute, and a trustee of the Samuel H. Kress Foundation and the Public Broadcasting Service.

STEPHEN M. PECK was elected to our supervisory board in 1999. He is currently a partner with Wilderness Funds LP. He is a former managing partner of Weiss, Peck & Greer which he co-founded in 1970. He served as Chief Investment Officer and Director of Reliance Insurance Company, Inc. from January 1986 to July 1988. Mr. Peck is a member of the Board of Directors of Advance Auto Parts, Inc., Boston Life Sciences, Inc.

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and Carnac Resource, Inc. Mr. Peck is presently on the Board of Trustees of Mount Sinai Medical Center, Mount Sinai Hospital, and the Mount Sinai School of Medicine. He is a member of the Board of Trustees of Mount Sinai/ NYU Health and The Jewish Theological Seminary.

Management Board

Each member of our management board is appointed by the supervisory board for a maximum term of five years and is eligible for reappointment thereafter. Their terms expire at our annual general meeting in the years listed below.

The table below provides names, ages, positions and terms of office of the members of our management board as of December 31, 2003.

Name	Age as of December 31, 2003	Position	Year Term Expires
Dr. Ben J. Lipps	63	Chairman of the management board, Chief Executive Officer of our Company and Chief Executive Officer for North America	2008
Roberto Fusté	51	Chief Executive Officer for Asia Pacific	2006
Dr. Emanuele Gatti	48	Chief Executive Officer for Europe, Middle East, Africa and Latin America	2005
Lawrence Rosen	45	Chief Financial Officer (as of November 1, 2003)	2005
Dr. Rainer Runte	44	General Counsel and Chief Compliance Officer	2005

DR. BEN J. LIPPS has been Chairman of the management board since May 1, 1999 and was Vice Chairman of the management board from September 1998 until May 1, 1999. He has been President and a director of Fresenius Medical Care Holdings since September 1996 and President, Chief Executive Officer, Chief Operating Officer and a director of Fresenius USA since October 1989, and served in various capacities with Fresenius USA's predecessor since 1985. Dr. Lipps joined Dow Chemical Company in 1966 and led the research team that developed the first hollow fiber dialyzer between 1967 and 1969. Prior to joining Fresenius USA's predecessor, Dr. Lipps was a Vice President of Research and Development for Cordis Dow Corporation.

ROBERTO FUSTÉ was appointed to our management board effective January 1, 1999. Mr. Fusté is responsible for the Asia-Pacific region within the International segment, for which he assumed responsibility in 1998. Mr. Fusté joined Fresenius AG in 1991 when Fresenius AG acquired Nephrocontrol S.A., a Spanish company which he founded in 1985 and of which he was Managing Director and joint owner. After the company was acquired by Fresenius AG, he continued as Managing Director. In 1995, he joined the Head Office of Fresenius AG where he has held various executive positions.

DR. EMANUELE GATTI has been a member of our management board since May 1997 and is President and Chief Executive Officer of Europe, Latin America, Middle East and Africa within the International segment. Previously he was Executive Vice President with responsibility for our dialysis business in Southern Europe. Dr. Gatti joined the Fresenius Group in 1989 when Fresenius AG acquired Sis-ter, an Italian company of which he was General Manager. He has been working in the field of dialysis since 1981 after leaving the Polytechnic School of Milan where he was involved in teaching and biomedical research.

LAWRENCE ROSEN joined our management board on November 1, 2003 and is Chief Financial Officer. He served as Group Senior Vice President for Corporate Finance and Treasury with Aventis S.A., in Strasbourg, France before his move to Fresenius Medical Care. In that position he was responsible for all Corporate Finance activities as well as Risk and Cash Management, including banking and rating agency relationships. Lawrence Rosen worked for Aventis S.A. and its predecessor companies since 1984 and has served in senior finance and

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treasury positions in North America, Germany, and France. He holds a Master of Business Administration (MBA) from the University of Michigan, USA and a Bachelor of Science in Economics from the State University of New York at Brockport, USA.

DR. RAINER RUNTE was appointed a deputy member of our management board as General Counsel and Chief Compliance Officer effective March 15, 2002 and as of January 1, 2004 was made a full member of the management board responsible for law and compliance worldwide. Dr. Runte has worked as in-house counsel within Fresenius AG and its subsidiaries and has served as General Counsel of Fresenius Medical Care since 1997.

On February 25, 2004, the following new members were appointed to the Company's management board:

RICE POWELL age 48, has more than 25 years of experience in the healthcare industry. Since 1997 he has been the President of Renal Products division of Fresenius Medical Care in North America including the Extracorporeal Therapy and Laboratory Services.

MATS WAHLSTROM, age 49, also has nearly 20 years of experience in the renal field. Since November 2002 he has been the President of Fresenius Medical Care's Medical Services division in North America.

The business address of all members of our management board and supervisory board is Else-Kröner-Strasse 1, 61352 Bad Homburg, Germany.

B. Compensation

Compensation of Our Management Board and our Supervisory Board

For the year ended December 31, 2003, we paid aggregate cash compensation to all members of the management board of 3.4 million. The aggregate compensation fees to all members of the supervisory board was 0.4 million including compensation to Dr. Krick for his duties as Chairman of the supervisory board. We pay an annual retainer fee to each member of the supervisory board, with the Chairman paid twice that amount and the Deputy Chairman paid 150% of that amount. We reimburse supervisory board members for their reasonable travel and accommodation expenses incurred with respect to their duties as supervisory board members. The aggregate compensation reported above does not include amounts paid as fees for services rendered by certain business or professional entities with which some of the supervisory board members are associated.

During 2003 we awarded no options to members of the management board to purchase our preference shares without or without stock price targets under the new FMC International 2001 Plan. At December 31, 2003 management board members held options to acquire 99,600 Preference shares, all of which were exercisable at a weighted average exercise price of 36.49 under FMC 98 Plan 2 and 239,250 options, of which 27,460 are exercisable under the FMC 2001 stock incentive plan.

During 1999, the Company granted to a member of the management board a five-year loan of \$2 million with interest at 6.0% per annum. This loan was repaid in 2003.

C. Board Practices

For information relating to the terms of office of our management board and our supervisory board and the periods in which the members of those bodies have served in office, see Item 6.A. above. We do not have a remuneration committee. Our supervisory board carries out the functions usually performed by the remuneration committee, and our supervisory board reviews the compensation of the members of our management board. On March 6, 2003 the supervisory board created an audit committee whose members are Dr. Bernd Fahrholz, Walter Weisman and Stephen M. Peck. The primary function of the audit committee is to assist the Board in fulfilling its oversight responsibilities, primarily through:

overseeing management's conduct or our financial reporting process and the internal accounting and financial control systems and auditing of our financial statements;

monitoring our internal controls risk program;

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monitoring the independence and performance of our outside auditors;

providing an avenue of communication among the outside auditors, management and the supervisory board;

retaining the services of our independent auditors (subject to the approval by our shareholders at our annual general meeting) and approval of their fees; and

pre-approval of all audit and non-audit services performed by KPMG Deutsche Treuhand-Gesellschaft AG Wirtschaftsprüfungsgesellschaft, the accounting firm which audits our consolidated financial statements.

D. Employees

At December 31, 2003, we had 41,097 employees, as compared to 39,264 at December 31, 2002 and 37,331 at December 31, 2001. They are employed in our principal segments as follows: North America 26,953 employees and International 14,144. The following table shows the average number of employees by segment and our major category of activities for the last three fiscal years.

	<u>2003</u>	<u>2002</u>	<u>2001</u>
North America			
Dialysis Care	21,986	21,628	22,382
Dialysis Products	4,967	4,861	3,970
	<u>26,953</u>	<u>26,489</u>	<u>26,352</u>
International			
Dialysis Care	7,788	6,924	6,173
Dialysis Products	6,356	5,851	4,806
	<u>14,144</u>	<u>12,775</u>	<u>10,979</u>

We are a member of the Chemical Industry Employers Association in Germany and we are bound by union agreements negotiated with the respective union representatives. We are also party to additional shop agreements negotiated with works councils at individual facilities that relate to those facilities. In addition, approximately 2% of our U.S. employees are covered by collective bargaining agreements. During the last three fiscal years, we have not suffered any labor-related work disruptions.

E. Share ownership

As of December 31, 2003, members of the supervisory board and the management board as a group owned 4,367 Ordinary shares (0.01% of total Ordinary shares outstanding) and 3,200 Preference shares (0.01% of total Preference shares outstanding). At December 31, 2003 management board members held options to acquire 338,850 Preference shares of which options to purchase 127,060 Preference shares were exercisable at a weighted average exercise price of 42.19. Those options expire at various dates between 2008 and 2013. None of the members of our management board and our supervisory board beneficially owns more than 1% of our outstanding Ordinary shares or our outstanding Preference shares.

Options to Purchase Our Securities**Stock Option Plans**

Immediately prior to the formation of Fresenius Medical Care, we adopted a stock incentive plan (the FMC Plan) for our key management and executive employees. As of December 31, 2003, 53,389 preference shares were available and exercisable with an average price range between \$55.59 and \$78.33 per share. Effective September 2001, no additional awards are granted under the FMC Plan.

During 1998, we adopted two stock incentive plans (FMC 98 Plan 1 and FMC 98 Plan 2) for our key management and executive employees. Under FMC 98 Plan 1, eligible employees have the right to acquire Preference shares of the Company. The maximum number of Preference shares that may be issued under this plan

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is 2,443,333 less any shares issued, or subject to issue, under the FMC Plan. Any shares available due to forfeiture of Grants under the FMC Plan would be considered available under FMC 98 Plan 1 as long as the total Preference shares issued under both plans does not exceed the 2,443,333 shares noted above. Under FMC 98 Plan 2, eligible employees have the right to acquire our Preference shares (Options). The share price of the Preference share shall be equal to the average of the official daily quotation prices of the Preference shares on the Frankfurt Stock Exchange on the thirty days (30) of trading immediately prior to the date of grant of the Option. One-third of an Option vests on each of the second, third and fourth anniversaries of the award date, provided that we achieve certain performance criteria for the full fiscal year following the grant date in comparison to the full fiscal year preceding the grant date. On May 30, 2000, our shareholders approved a change to the FMC 98 Plan 2 whereby the impact of the special charge for the 1999 Settlement was excluded from the our performance criteria relative to the EBIT growth requirements in the plan. Options granted under FMC 98 Plan 2 have a 10-year term. The maximum number of Preference shares that may be issued under this plan is 2,500,000 shares, of which 500,000 are designated for management board members and 2,000,000 are for other managerial staff. Each option is exercisable into one Preference share. Effective September 2001, no additional Grants or Options will be awarded under FMC Plan 98 1 or FMC Plan 98 2.

On May 23, 2001, by resolution of our annual general meeting, the FMC 98 Plans were replaced by a new plan. The management board was empowered to issue convertible bonds with a total value of 10,240,000 to the members of the management board and to other employees entitling the holders to a total subscription of up to 4 million non-voting Preference shares. The convertible bonds have a par value of 2.56 and are interest bearing at a rate of 5.5%. Purchase of the bonds may be funded by a non-recourse loan secured by the bond with respect to which the loan was made. We have the right to offset our obligation on a convertible bond against the employee obligation on the related loan; therefore, the convertible bond obligations and employee loan receivables are not reflected in our consolidated financial statements. The bonds mature in ten years and are generally fully convertible after three years. The bonds may be issued either as convertible bonds which are subject to a stock price target or convertible bonds without a stock price target. In the case of convertible bonds which are subject to a stock price target the conversion right is exercisable only if the market price of the Preference shares increases by 25% or more over the grant-date price subsequent to the day of grant for at least one day prior to exercise. Participants have the right to opt for convertible bonds with or without the stock price target. In order to create an incentive to select convertible bonds which depend on the stock price target, the number of convertible bonds awarded to those employees who select the bonds without a stock price target will be reduced by 15%. Each convertible bond entitles the holder thereof, upon payment of a conversion price to convert the bond into one Preference share. The conversion price of the convertible bonds which are not subject to the stock price target is determined by the average price of the Preference shares during the last 30 trading days prior to the date of grant. The conversion price of the convertible bonds which depend on the stock price target corresponds to the closing price of the Preference shares the day the target was reached.

The managing board and supervisory board are authorized to issue up to 20% of the total number of convertible bonds each year through May 22, 2006. The plan is valid until the last convertible bond issued under this plan is terminated or converted. Of the 2,002,911 options outstanding at December 31, 2003, 360,049 options were issued under the plan with the stock price target.

Rollover options

In connection with our formation, employees of National Medical Care exchanged options to purchase W.R. Grace common stock and Fresenius USA employees exchanged options to purchase Fresenius USA common stock for equivalent options with respect to our Ordinary shares. When we were formed, German corporate law did not allow us to reserve Ordinary shares and issue them upon the exercise of these rollover options, as is done by U.S. corporations. Instead, we issued the Ordinary shares issuable upon exercise of the options to Fresenius AG, which will hold the shares until exercise of the options. Fresenius AG has agreed that it will not exercise voting power, and will return any dividends paid, with respect to the Ordinary shares underlying options formerly related to W.R. Grace common stock. Upon exercise of any of these options, the holder will pay the option exercise price to us and Fresenius AG will deliver the Ordinary shares to the depository for the Ordinary shares against issuance of American Depositary Shares (ADSs) representing Ordinary shares in the

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name of the option holder. Upon cancellation or expiration without exercise of options formerly relating to W.R. Grace common stock, Fresenius AG will transfer the underlying Ordinary shares to us at no cost. Upon cancellation or expiration without exercise of options formerly relating to Fresenius USA common stock, the underlying Ordinary shares will revert to Fresenius AG. All rollover options expire on the same date on which the previous options to purchase either the W.R. Grace common stock or Fresenius USA common stock to which such rollover options relate would have expired.

During the year ended December 31, 2003, 79,491 FMC Rollover Plan options were exercised by employees. Rollover Plan options for 24,927 Ordinary ADSs were outstanding and exercisable as of December 31, 2003 at a weighted average exercise price of \$13.53. Members of our supervisory board and our management board, as a group, held no rollover options.

Item 7. Major Shareholders and Related Party Transactions

A. Major Shareholders

Security Ownership of Certain Beneficial Owners of Fresenius Medical Care

Our outstanding share capital consists of Ordinary shares and non-voting Preference shares that are issued only in bearer form. Accordingly, unless we receive information regarding acquisitions of our shares through a filing with the Securities and Exchange Commission or through the German statutory requirements referred to below, we have no way of determining who our shareholders are or how many shares any particular shareholder owns except as described below with respect to our shares held in American Depository Receipt (ADR) form. Because we are a foreign private issuer under the rules of the Securities and Exchange Commission, our directors and officers are not required to report their ownership of our equity securities or their transactions in our equity securities pursuant to Section 16 of the Exchange Act. Under the German Securities Exchange Law (*Wertpapierhandelsgesetz*), holders of voting securities of a German company listed on the official market (*amtlicher Handel*) of a German stock exchange or a corresponding trading segment of a stock exchange within the European Union are obligated to notify the company of the level of their holding whenever such holding reaches, exceeds or falls below certain thresholds, which have been set at 5%, 10%, 25%, 50% and 75% of a company's outstanding voting rights.

We have been informed that as of December 31, 2003, Fresenius AG owned the majority, 50.8%, of our Ordinary shares. At December 31, 2003 Fresenius AG's Ordinary shares represented approximately 37% of our total share capital. JPMorgan Chase Bank, our ADR depository, informed us, that as of December 31, 2003 2,079,369 Ordinary ADSs, each representing one-third of an Ordinary share, were held of record by 7,673 U.S. holders and 47,401 Preference ADSs, each representing one-third of a Preference share, were held of record by 5 U.S. holders. Ordinary shares and Preference shares held directly by U.S. holders accounted for approximately 1% of our Ordinary shares outstanding and less than 1% of our Preference shares outstanding as of December 31, 2003. For more information regarding ADRs and ADSs see Item 10.B. Memorandum and Articles of Association Description of American Depositary Receipts.

According to a Schedule 13G filed with the Securities and Exchange Commission on February 17, 2004, by FMR Corp., Edward C. Johnson 3d and Abigail Johnson, at December 31, 2003, Edward C. Johnson 3d and Abigail Johnson were the beneficial owners of 3,649,110 or 5.2% of the common stock of Fresenius Medical Care AG. The Schedule 13G states that 2,458,237 such shares, or 3.5% of our common stock is beneficially owned by Fidelity Management & Research Company (Fidelity), 82 Devonshire Street, Boston, MA 02109, a wholly-owned subsidiary of FMR Corp. and a registered investment advisor to various investment companies registered under Section 8 of the Investment Company Act of 1940.

Security Ownership of Certain Beneficial Owners of Fresenius AG

Fresenius AG's share capital consists of Ordinary shares and non-voting Preference shares. Both classes of shares are issued only in bearer form. Accordingly, Fresenius AG has no way of determining who its shareholders are or how many shares any particular shareholder owns. However, under the German Securities Exchange Law, holders of voting securities of a German company listed on the official market (*amtlicher Handel*) of a German

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stock exchange or a corresponding trading segment of a stock exchange within the European Union are obligated to notify the company of certain levels of holdings, as described above.

Based on the most recent information available, Vermögensverwaltungsgesellschaft Nachlass Else Kröner mbH owns 67.4% of the Fresenius AG Ordinary shares. In addition, Allianz Lebensversicherungs-AG informed Fresenius AG that it owns 9.7% of the Fresenius AG Ordinary shares.

B. Related party transactions

In connection with the formation of Fresenius Medical Care, and the combination of the dialysis businesses of Fresenius AG and W.R. Grace, Fresenius AG and its affiliates and Fresenius Medical Care and its affiliates entered into several agreements for the purpose of giving effect to the merger and defining our ongoing relationship. Fresenius AG and W.R. Grace negotiated these agreements. The information below summarizes the material aspects of certain agreements, arrangements and transactions between Fresenius Medical Care and Fresenius AG and their affiliates. Some of these agreements have been previously filed with the Securities and Exchange Commission. The following descriptions are not complete and are qualified in their entirety by reference to the agreements, copies of which have been filed with the Securities and Exchange Commission and the New York Stock Exchange. We believe that the leases, the supply agreements and the service agreements are no less favorable to us and no more favorable to Fresenius AG than would have been obtained in arm's-length bargaining between independent parties. The trademark and other intellectual property agreements summarized below were negotiated by Fresenius AG and W.R. Grace, and, taken independently, are not necessarily indicative of market terms.

In the discussion below regarding our contractual and other relationships with Fresenius AG:

the term *we* (or *us*) and our affiliates refers *only* to Fresenius Medical Care AG and its subsidiaries; and

the term *Fresenius AG* and its affiliates refers *only* to Fresenius AG and affiliates of Fresenius AG *other than* Fresenius Medical Care AG and its subsidiaries.

Real Property Lease

We did not acquire the land and buildings in Germany that Fresenius Worldwide Dialysis used when we were formed. Fresenius AG or its affiliates have leased part of the real property to us, directly, and transferred the remainder of that real property to two limited partnerships. Fresenius AG is the sole limited partner of each partnership, and the sole shareholder of the general partner of each partnership. These limited partnerships, as landlords, have leased the properties to us and to Fresenius AG, as applicable, for use in our respective businesses. The aggregate annual rent payable by us under these leases is approximately 11.8 million, which was approximately \$13.3 million as of December 31, 2003, exclusive of maintenance and other costs, and is subject to escalation, based upon the German cost of living index for a four-person employee household. The leases for manufacturing facilities have a ten-year term, followed by two successive optional renewal terms of ten years each at our election. The leases for the other facilities have a term of ten years. Based upon an appraisal, we believe that the rents under the leases represent fair market value for such properties. For information with respect to our principal properties in Germany, see Item 4.D. Property, plants and equipment.

Covenants Not to Compete

Each of Fresenius AG and W.R. Grace has agreed that, for a period of ten years after our formation, it will not compete with us in any aspect of the business of supplying renal care-related goods and services, including laboratories. However, Fresenius AG may continue its home care business.

Trademarks

Fresenius AG continues to own the name and mark *Fresenius* and its *F* logo. Fresenius AG and Fresenius Medical Care Deutschland GmbH, our principal German subsidiary, have entered into agreements containing the following provisions. Fresenius AG has granted to our German subsidiary, for our benefit and that

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of our affiliates, an exclusive, worldwide, royalty-free, perpetual license to use Fresenius Medical Care in our corporate names, and to use the Fresenius marks, including some combination marks containing the Fresenius name that were used by Fresenius AG's dialysis business, and the Fresenius Medical Care name as a trade name, in all aspects of the renal business. Our German subsidiary, for our benefit and that of our affiliates, has also been granted a worldwide, royalty-free, perpetual license:

to use the Fresenius Medical Care mark in the then current National Medical Care non-renal business if it is used as part of Fresenius Medical Care together with one or more descriptive words, such as Fresenius Medical Care Home Care or Fresenius Medical Care Diagnostics ;

to use the F logo mark in the National Medical Care non-renal business, with the consent of Fresenius AG. That consent will not be unreasonably withheld if the mark using the logo includes one or more additional descriptive words or symbols; and

to use Fresenius Medical Care as a trade name in both the renal business and the National Medical Care non-renal business.

We and our affiliates have the right to use Fresenius Medical Care as a trade name in other medical businesses only with the consent of Fresenius AG. Fresenius AG may not unreasonably withhold its consent. In the U.S. and Canada, Fresenius AG will not use Fresenius or the F logo as a trademark or service mark, except that it is permitted to use Fresenius in combination with one or more additional words such as Pharma Home Care as a service mark in connection with its home care business and may use the F logo as a service mark with the consent of our principal German subsidiary. Our subsidiary will not unreasonably withhold its consent if the service mark includes one or more additional descriptive words or symbols. Similarly, in the U.S. and Canada, Fresenius AG has the right to use Fresenius as a trade name, but not as a mark, only in connection with its home care and other medical businesses other than the renal business and only in combination with one or more other descriptive words, provided that the name used by Fresenius AG is not confusingly similar to our marks and trade names. After the expiration of Fresenius AG's ten-year covenant not to compete with us, Fresenius AG may use Fresenius in its corporate names if it is used in combination with one or more additional descriptive word or words, provided that the name used by Fresenius AG is not confusingly similar to the Fresenius Medical Care marks or corporate or trade names.

Other Intellectual Property

Some of the patents, patent applications, inventions, know-how and trade secrets that Fresenius Worldwide Dialysis used prior to our formation were also used by other divisions of Fresenius AG. For Biofine, the polyvinyl chloride-free packaging material, Fresenius AG has granted to our principal German subsidiary, for our benefit and for the benefit of our affiliates, an exclusive license for the renal business and a non-exclusive license for all other fields except other non-renal medical businesses. Our German subsidiary and Fresenius AG will share equally any royalties from licenses of the Biofine intellectual property by either our German subsidiary or by Fresenius AG to third parties outside the renal business and the other non-renal medical businesses. In addition, Fresenius AG has transferred to our German subsidiary the other patents, patent applications, inventions, know-how and trade secrets that were used predominantly in Fresenius AG's dialysis business. In certain cases Fresenius Worldwide Dialysis and the other Fresenius AG divisions as a whole each paid a significant part of the development costs for patents, patent applications, inventions, know-how and trade secrets that were used by both prior to the merger. Where our German subsidiary acquired those jointly funded patents, patent applications, inventions, know-how and trade secrets, our subsidiary licensed them back to Fresenius AG exclusively in the other non-renal medical businesses and non-exclusively in all other fields. Where Fresenius AG retained the jointly funded patents, patent applications, inventions, know-how and trade secrets, Fresenius AG licensed them to our German subsidiary exclusively in the renal business and non-exclusively in all other fields.

Supply Agreements

We produce most of our products in our own facilities. However, Fresenius AG manufactures some of our products for us, principally dialysis concentrates, at facilities that Fresenius AG retained. These facilities are

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located in Brazil and France. Conversely, a facility in Italy that Fresenius AG transferred to us produces products for Fresenius Kabi AG, a subsidiary of Fresenius AG.

Our local subsidiaries and those of Fresenius AG have entered into supply agreements for the purchase and sale of products from the above facilities. Prices under the supply agreements include a unit cost component for each product and an annual fixed cost charge for each facility. The unit cost component, which is subject to annual review by the parties, is intended to compensate the supplier for variable costs such as costs of materials, variable labor and utilities. The fixed cost component generally will be based on an allocation of the 1995 fixed costs of each facility, such as rent, depreciation, production scheduling and quality control. The fixed cost component will be subject to adjustment by good-faith negotiation every twenty-four months. If the parties cannot agree upon an appropriate adjustment, the adjustment will be made based on an appropriate consumer price index in the country in which the facility is located.

Each supply agreement has a term that is approximately equal to the estimated average life of the relevant production assets, resulting in terms of four and one-half to five years. Each supply agreement may be terminated by the purchasing party after specified notice period, subject to a compensation payment reflecting a portion of the relevant fixed costs.

The parties may modify existing or enter into additional supply agreements, arrangements and transactions. Any future modifications, agreements, arrangements and transactions will be negotiated between the parties and will be subject to the approval provisions of the pooling agreements and the regulatory provisions of German law regarding dominating enterprises.

Services Agreement

We obtain administrative and other services from Fresenius AG headquarters and from other divisions and subsidiaries of Fresenius AG. These services relate to, among other things, data processing, financial and management accounting and audit, human resources, risk management, quality control, production management, research and development, marketing and logistics. For 2003, Fresenius AG charged us approximately \$26.6 million for these services. Conversely, we have provided certain services to other divisions and subsidiaries of Fresenius AG relating to research and development, plant administration, patent administration and warehousing. For 2003, we charged approximately \$11.7 million to Fresenius AG's other divisions and subsidiaries for services we rendered to them.

We and Fresenius AG may modify existing or enter into additional services agreements, arrangements and transactions. Any such future modifications, agreements, arrangements and transactions will be negotiated between the parties and will be subject to the approval provisions of the pooling agreements and the regulations of German law regarding dominating enterprises.

Financing

At December 31, 2003, aggregate loans outstanding from Fresenius AG amounted to \$30 million which bore interest at 1.165% at year end. The borrowed funds were used for general corporate purposes. Interest paid during 2003 was \$0.1 million.

Purchase of the Adsorber Business

In 2003, we purchased the adsorber business of Fresenius AG for a purchase price of \$27.3 million, net of cash acquired. The adsorber business manufactures products used in the field of therapeutic apheresis. These therapies are similar to kidney dialysis treatment in that they consist of extracorporeal blood treatment. In accordance with the requirements of the pooling agreements relating to our outstanding Ordinary shares and Preferred shares, the acquisition of the Fresenius AG adsorber business was approved by our independent directors.

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Other Interests

During 1999, we granted a member of our management board a five-year unsecured loan in the amount of \$2 million, with interest at 6.0% per annum. This loan was repaid in 2003.

Prof. Dr. Bernd Fahrholz, a member of our supervisory board, was the chairman of the management board of Dresdner Bank AG until 2003 and Dr. Manfred Schaudwet, the retired Executive Manager of Dresdner Bank AG, is a member of the supervisory board of Fresenius AG. See Security Ownership of Certain Beneficial Owners of Fresenius AG. Dresdner Bank AG, through its New York and Cayman branches, is a documentation agent and one of the joint lead arrangers and book managers under 2003 Senior Credit Agreement. It was also one of four co-arrangers of our prior principal credit agreement and one of the managing agents under that facility. Dresdner Bank AG also acts as custodian under the deposit agreement for the ADSs evidencing our Ordinary shares and under the deposit agreement for the ADSs evidencing our Preference shares, and an affiliate of Dresdner Kleinwort Wasserstein Securities LLC (a wholly-owned subsidiary of Dresdner Bank AG) is the New York Stock Exchange specialist for the ADSs evidencing our Ordinary shares.

Dr. Dieter Schenk, Deputy Chairman of our supervisory board and a member of the supervisory board of Fresenius AG, is a partner in the law firm of Nörr Stiefenhofer Lutz, which has provided legal services to Fresenius AG and Fresenius Medical Care. See Security Ownership of Certain Beneficial Owners of Fresenius AG. Dr. Schenk is one of the executors of the estate of Mrs. Else Kröner. Vermögensverwaltungsgesellschaft Nachlass Else Kröner mbH, a charitable foundation established under the will of Mrs. Kröner, owns the majority of the voting shares of Fresenius AG.

Products

During 2003, we recognized \$27.3 million of sales to Fresenius AG and its affiliates. We made purchases from Fresenius AG in the amount of \$27.2 million during 2003.

Item 8. Financial information

The information called for by parts 8.A.1 through 8.A.6 of this item is in the section beginning on Page F-1.

8.A.7. Legal Proceedings

This section describes material legal actions and proceedings relating to us and our business.

Commercial Litigation

We were formed as a result of a series of transactions pursuant to the Agreement and Plan of Reorganization (the Merger) dated as of February 4, 1996 by and between W.R. Grace & Co. and Fresenius AG. At the time of the Merger, a W.R. Grace & Co. subsidiary known as W.R. Grace & Co.-Conn. had, and continues to have, significant potential liabilities arising out of product-liability related litigation, pre-Merger tax claims and other claims unrelated to NMC, which was Grace's dialysis business prior to the Merger. In connection with the Merger, W.R. Grace & Co.-Conn. agreed to indemnify us, FMCH, and NMC against all liabilities of W.R. Grace & Co., whether relating to events occurring before or after the Merger, other than liabilities arising from or relating to NMC's operations. W.R. Grace & Co. and certain of its subsidiaries filed for reorganization under Chapter 11 of the U.S. Bankruptcy Code (the Grace Chapter 11 Proceedings) on April 2, 2001.

Pre-Merger tax claims or tax claims that would arise if events were to violate the tax-free nature of the Merger, could ultimately be our obligation. In particular, W. R. Grace & Co. has disclosed in its filings with the Securities and Exchange Commission that: its tax returns for the 1993 to 1996 tax years are under audit by the Internal Revenue Service (the Service); W.R. Grace & Co. has received the Service's examination report on tax periods 1993 to 1996; that during those years Grace deducted approximately \$122 million in interest attributable to corporate owned life insurance (COLI) policy loans; that W.R. Grace & Co. has paid \$21 million of tax and interest related to COLI deductions taken in tax years prior to 1993; that a U.S. District Court ruling has denied interest deductions of a taxpayer in a similar situation and that W.R. Grace & Co. is seeking a settlement of the Service's claims. Subject to certain representations made by W.R. Grace & Co.,

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Fresenius Medical Care AG and Fresenius AG, W.R. Grace & Co. and certain of its affiliates agreed to indemnify us against this and other pre-Merger and Merger related tax liabilities.

Prior to and after the commencement of the Grace Chapter 11 Proceedings, class action complaints were filed against W.R. Grace & Co. and FMCH by plaintiffs claiming to be creditors of W.R. Grace & Co.-Conn., and by the asbestos creditors committees on behalf of the W.R. Grace & Co. bankruptcy estate in the Grace Chapter 11 Proceedings, alleging among other things that the Merger was a fraudulent conveyance, violated the uniform fraudulent transfer act and constituted a conspiracy. All such cases have been stayed and transferred to or are pending before the U.S. District Court as part of the Grace Chapter 11 Proceedings.

In 2003, we reached an agreement with the asbestos creditors committees on behalf of the W.R. Grace & Co. bankruptcy estate and W.R. Grace & Co. in the matters pending in the Grace Chapter 11 Proceedings for the settlement of all fraudulent conveyance and tax claims against it and other claims related to us that arise out of the bankruptcy of W.R. Grace & Co. Under the terms of the settlement agreement as amended (the Settlement Agreement), fraudulent conveyance and other claims raised on behalf of asbestos claimants will be dismissed with prejudice and we will receive protection against existing and potential future W.R. Grace & Co. related claims, including fraudulent conveyance and asbestos claims, and indemnification against income tax claims related to the non-NMC members of the W.R. Grace & Co. consolidated tax group upon confirmation of a W.R. Grace & Co. bankruptcy reorganization plan that contains such provisions. Under the Settlement Agreement, we will pay a total of \$115 million to the W.R. Grace & Co. bankruptcy estate, or as otherwise directed by the Court, upon plan confirmation. No admission of liability has been or will be made. The Settlement Agreement has been approved by the U.S. District Court. Subsequent to the Merger, W.R. Grace & Co. was involved in a multi-step transaction involving Sealed Air Corporation (formerly known as Grace Holding, Inc.). We are engaged in litigation with Sealed Air Corporation (Sealed Air) to confirm our entitlement to indemnification from Sealed Air for all losses and expenses incurred by us relating to pre-Merger tax liabilities and Merger-related claims. Under the Settlement Agreement, upon confirmation of a plan that satisfies the conditions of our payment obligation, this litigation will be dismissed with prejudice.

On April 4, 2003, FMCH filed a suit in the United States District Court for the Northern District of California, *Fresenius USA, Inc., et al., v. Baxter International Inc., et al.*, Case No. C 03-1431, seeking a declaratory judgment that we do not infringe on patents held by Baxter International Inc. and its subsidiaries and affiliates (Baxter), that the patents are invalid, and that Baxter is without right or authority to threaten or maintain suit against us for alleged infringement of Baxter's patents. In general, the alleged patents concern touch screens, conductivity alarms, power failure data storage, and balance chambers for hemodialysis machines. Baxter has filed counterclaims against us seeking monetary damages and injunctive relief, and alleging that we willfully infringed on Baxter's patents. We believe our claims are meritorious, although the ultimate outcome of any such proceedings cannot be predicted at this time and an adverse result could have a material adverse effect on our business, financial condition, and results of operations.

Other Litigation and Potential Exposures

From time to time, we are a party to or may be threatened with other litigation arising in the ordinary course of our business. Management regularly analyzes current information including, as applicable, our defenses and insurance coverage and, as necessary, provides accruals for probable liabilities for the eventual disposition of these matters.

We, like other health care providers, conduct our operations under intense government regulation and scrutiny. We must comply with regulations which relate to or govern the safety and efficacy of medical products and supplies, the operation of manufacturing facilities, laboratories and dialysis clinics, and environmental and occupational health and safety. We must also comply with the Anti-Kickback Statute, the False Claims Act, the Stark Statute, and other federal and state fraud and abuse laws. Applicable laws or regulations may be amended, or enforcement agencies or courts may make interpretations that differ from ours or the manner in which we conduct our business. In the U.S. enforcement has become a high priority for the federal government and some states. In addition, the provisions of the False Claims Act authorizing payment of a portion of any recovery to the party bringing the suit encourage private plaintiffs to commence whistle blower actions. By virtue of this

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regulatory environment, as well as our corporate integrity agreement with the government, we expect that our business activities and practices will continue to be subject to extensive review by regulatory authorities and private parties, and expect continuing inquiries, claims and litigation relating to our compliance with applicable laws and regulations. We may not always be aware that an inquiry or action has begun, particularly in the case of whistle blower actions, which are initially filed under court seal.

We operate many facilities throughout the U.S. In such a decentralized system, it is often difficult to maintain the desired level of oversight and control over the thousands of individuals employed by many affiliated companies. We rely upon our management structure, regulatory and legal resources, and the effective operation of our compliance program to direct, manage and monitor the activities of these employees. On occasion, we may identify instances where employees, deliberately or inadvertently, have submitted inadequate or false billings. The actions of such persons may subject us and our subsidiaries to liability under the Anti-Kickback Statute, the Stark Statute and the False Claims Act, among other laws.

Physicians, hospitals and other participants in the health care industry are also subject to a large number of lawsuits alleging professional negligence, malpractice, product liability, worker's compensation or related claims, many of which involve large claims and significant defense costs. We have been subject to these suits due to the nature of our business and we expect that those types of lawsuits may continue. Although we maintain insurance at a level which we believe to be prudent, we cannot assure that the coverage limits will be adequate or that insurance will cover all asserted claims. A successful claim against us or any of our subsidiaries in excess of insurance coverage could have a material adverse effect upon us and the results of our operations. Any claims, regardless of their merit or eventual outcome, also may have a material adverse effect on our reputation and business.

We have also had claims asserted against us and had lawsuits filed against us relating to businesses that we have acquired or divested. These claims and suits relate both to operation of the businesses and to the acquisition and divestiture transactions. We have asserted our own claims, and claims for indemnification. Although the ultimate outcome cannot be predicted at this time, an adverse result could have a material adverse effect upon our business, financial condition, and results of operations.

Accrued Special Charge for Legal Matters

At December 31, 2001, we recorded a pre-tax special charge of \$258 million to reflect anticipated expenses associated with the defense and resolution of pre-Merger tax claims, Merger-related claims, and commercial insurer claims. The costs associated with the Settlement Agreement and settlements with insurers have been charged against this accrual. While we believe that our remaining accruals reasonably estimate our currently anticipated costs related to the continued defense and resolution of the remaining matters, no assurances can be given that our actual costs incurred will not exceed the amount of this accrual.

8.A.8. Dividend Policy

We generally pay annual dividends on both our Preference shares and our Ordinary shares in amounts that we determine on the basis of the prior year unconsolidated earnings of Fresenius Medical Care AG as shown in the statutory financial statements that we prepare under German law, subject to authorization by a resolution to be passed at our general meeting of shareholders. Under our articles of association, the minimum dividend payable on the Preference shares is 0.12 per share and, if we declare dividends, holders of our Preference shares must receive 0.06 per share more than the dividend on an Ordinary share. Under German law, we must, in all cases, pay the annual dividend declared on our Preference shares before we pay dividends declared on our Ordinary shares.

Our management board and our supervisory board propose dividends and the shareholders approve dividends for payment in respect of a fiscal year at the annual general meeting in the following year. Since all of our shares are in bearer form, we either remit dividends to the depository bank (*Depotbank*) on behalf of the shareholders or, in the case of shareholders holding physical certificates, we pay dividends through the paying agents that we appoint against presentation of the relevant dividend coupon. Details of the paying agents are published in the German Federal Gazette (*Bundesanzeiger*).

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Our senior credit agreement as well as the senior subordinated indentures relating to our trust preferred securities restrict our ability to pay dividends. Item 5.B. Operating and Financial Review and Prospects Liquidity and Capital Resources and the notes to our consolidated financial statements appearing elsewhere in this report discuss this restriction.

The table below provides information regarding the annual dividend per share that we paid on our Preference shares and Ordinary shares. The dividends shown for each year were paid with respect to our operations in the preceding year.

Per Share Amount	2003	2002	2001
Preference share	0.94	0.91	0.84
Ordinary share	1.00	0.85	0.78

We have announced that our managing board and our supervisory board have proposed dividends for 2003 payable in 2004 of 1.08 per Preference share and 1.02 per Ordinary share. These dividends are subject to approval by our shareholders at our annual general meeting to be held on May 27, 2004.

Except as described herein, holders of ADSs will be entitled to receive dividends on the Ordinary shares and the Preference shares represented by the respective ADSs. We will pay any cash dividends payable to such holders to the depositary in euros and, subject to certain exceptions, the depositary will convert the dividends into U.S. dollars. Fluctuations in the exchange rate between the U.S. dollar and the euro will affect the amount of dividends that ADS holders receive. Dividends paid on the Preference shares and dividends paid to holders and beneficial holders of the ADSs will be subject to deduction of German withholding tax. You can find a discussion of German withholding tax below in Item 10.E. Taxation .

B. Significant Changes

None

Item 9. The Offer and Listing Details**A.4. and C. Information regarding the trading markets for price history of our stock****Trading Markets**

The principal trading market for the Ordinary shares and the Preference shares is the Frankfurt Stock Exchange. All Ordinary shares and Preference shares have been issued in bearer form. Accordingly, we have no way of determining who our holders of Ordinary and Preference shares are or how many shares any particular shareholder owns, with the exception of the number of shares held in ADR form in the United States. For more information regarding ADRs see Item 10.B. Memorandum and articles of association Description of American Depositary Receipts. However, under the German Stock Corporation and Securities Law, holders of voting securities of a German company listed on a stock exchange within the EU are obligated to notify the company of certain levels of holdings as described in Item 7.A. Major Shareholders . The Ordinary shares have been listed on the Frankfurt Stock Exchange since October 2, 1996. The Preference shares have been listed on the Frankfurt Stock Exchange since November 25, 1996.

Since October 1, 1996, ADSs each representing one-third of an Ordinary share (the Ordinary ADSs), have been listed and traded on the New York Stock Exchange (NYSE) under the symbol FMS. Since November 25, 1996, ADSs, each representing one-third of a Preference share (the Preference ADSs), have been listed and traded on the NYSE under the symbol FMS- p. The Depositary for both the Ordinary ADSs and the Preference ADSs is Morgan Guaranty Trust Company of New York (the Depositary).

Trading on the Frankfurt Stock Exchange

Deutsche Börse AG operates the Frankfurt Stock Exchange, which is the most significant of the eight German stock exchanges. As of December 31, 2003, the most recent figures available, the shares of 5,730

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companies traded on the official market, regulated market and the regulated unofficial market of the Frankfurt Stock Exchange. Of these 820 were German companies and 4,901 were foreign companies.

Trading on the floor of the Frankfurt Stock Exchange begins every business day at 9:00 a.m. and ends at 8:00 p.m., Central European Time (CET). Securities listed on the Frankfurt Stock Exchange generally trade in the auction market, but also change hands in interbank dealer markets. Prices are noted by publicly commissioned stock brokers who are members of the Frankfurt Stock Exchange, but who do not as a rule deal with the public. These prices are determined by out-cry. The prices of actively traded securities, including the shares of large corporations, are continuously quoted during trading hours. For all securities, a fixed price (*Einheitspreis*) is established at approximately midday on each day the Frankfurt Stock Exchange is open for business.

FMC s shares are traded on Xetra (Exchange Electronic Trading) in addition to being traded on the auction market. Starting on November 3, 2003, the Deutsche Börse AG shortened the trading hours for Xetra to between 9:00 a.m. and 5:30 p.m. CET instead of between 9:00 a.m. and 8:00 p.m. These hours are effective for one year, and will be reviewed after 6 months to decide if they should be continued. Only brokers and banks that have been admitted to Xetra by the Frankfurt Stock Exchange may trade on the system. Private investors can trade on Xetra though their banks and brokers.

Deutsche Börse AG publishes an official daily list of quotations (*Amtliches Kursblatt*) containing the fixed prices (*Einheitskurse*) and other information for all traded securities on the Internet, webpage <http://www.exchange.de>.

Transactions on the Frankfurt Stock Exchange (including transactions through the Xetra system) settle on the second business day following the trade. Transactions off the Frankfurt Stock Exchange (such as, for example, large trades or transactions in which one of the parties is foreign) generally also settle on the second business day following the trade, although a different period may be agreed to by the parties. Under standard terms and conditions for securities transactions employed by German banks, customers orders for listed securities must be executed on a stock exchange unless the customer gives specific instructions to the contrary.

The Frankfurt Stock Exchange can suspend a quotation if orderly trading is temporarily endangered or if a suspension is deemed to be necessary to protect the public.

The Hessian Stock Exchange Supervisory Authority and the Trading Monitoring Unit of the Frankfurt Stock Exchange, which is under the control of the Stock Exchange Supervisory Authority, both monitor trading on the Frankfurt Stock Exchange.

The Federal Supervisory Authority for Securities Trading (*Bundesaufsichtsamt für den Wertpapierhandel*), an independent federal authority, is responsible for the general supervision of securities trading pursuant to provisions of the German Securities Trading Act (*Wertpapierhandelsgesetz*).

The table below sets forth for the periods indicated, the high and low closing sales prices in euro for the Ordinary shares on the Frankfurt Stock Exchange, as reported by the Frankfurt Stock Exchange Xetra system. Since January 4, 1999, all shares on German stock exchanges trade in euro. (See the discussion under Item 11. Quantitative and Qualitative Disclosures about Market Risk with respect to the rates of exchange between the Dollar, euro and deutsche mark).

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		Price per ordinary share ()	
		High	Low
2004	February	56.63	52.14
	January	58.03	52.16
2003	December	57.00	53.07
	November	54.99	49.06
	October	52.95	48.25
	September	53.77	45.61
2003	Fourth Quarter	57.00	48.25
	Third Quarter	53.77	42.00
	Second Quarter	50.90	39.32
	First Quarter	48.79	38.00
2002	Fourth Quarter	46.40	20.76
	Third Quarter	53.19	19.98
	Second Quarter	73.00	42.00
	First Quarter	71.21	56.00
2003	Annual	57.00	38.00
2002	Annual	73.00	19.98
2001	Annual	92.90	66.77
2000	Annual	103.60	72.40
1999	Annual	88.70	47.20

The average daily trading volume of the Ordinary shares traded on the Frankfurt Stock Exchange during 2003 was 340,579 shares. The foregoing numbers are based on total yearly turnover statistics supplied by the Frankfurt Stock Exchange. On February 27, 2004, the closing sales price per Ordinary share on the Frankfurt Stock Exchange was 53.94, equivalent to \$69.25 per Ordinary share.

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The table below sets forth for the periods indicated, the high and low closing sales prices in euro for the Preference shares on the Frankfurt Stock Exchange, as reported by the Frankfurt Stock Exchange. As all shares on German stock exchanges trade in euro since January 4, 1999 (see the discussion under Item 11. Quantitative and Qualitative Disclosures about Market Risk with respect to the rates of exchange between the Dollar, euro and deutsche mark).

		Price per preference share ()	
		High	Low
2004	February	38.33	36.20
	January	40.95	35.84
2003	December	41.00	38.25
	November	40.17	35.26
	October	39.50	35.01
	September	40.50	33.80
	August	34.58	31.40
2003	Fourth Quarter	41.00	35.01
	Third Quarter	40.50	30.09
	Second Quarter	36.00	28.50
	First Quarter	35.60	27.36
2002	Fourth Quarter	33.80	15.80
	Third Quarter	38.00	15.17
	Second Quarter	53.90	33.10
	First Quarter	53.80	41.50
2003	Annual	41.00	28.50
2002	Annual	53.90	15.17
2001	Annual	65.25	46.01
2000	Annual	58.00	38.00
1999	Annual	43.50	30.30

The average daily trading volume of the Preference shares traded on the Frankfurt Stock Exchange during 2003 was 34,456 shares. The foregoing numbers are based on total yearly turnover statistics supplied by the Frankfurt Stock Exchange. On February 27, 2004, the closing sales price per Preference share on the Frankfurt Stock Exchange was 36.49, equivalent to \$46.84 per Preference share.

Table of Contents**Trading on the New York Stock Exchange**

The table below sets forth, for the periods indicated, the high and low closing sales prices for the Ordinary ADSs on the NYSE:

		Price per ordinary ADS (\$)	
		High	Low
2004	February	23.90	21.90
	January	24.05	21.76
2003	December	23.54	21.53
	November	21.66	18.80
	October	20.39	18.90
	September	20.20	16.79
2003	Fourth Quarter	23.54	18.80
	Third Quarter	20.20	16.00
	Second Quarter	18.00	15.33
	First Quarter	17.49	13.20
2002	Fourth Quarter	15.20	7.04
	Third Quarter	18.10	6.70
	Second Quarter	21.60	14.20
	First Quarter	21.20	16.35
2003	Annual	23.54	13.20
2002	Annual	21.60	6.70
2001	Annual	28.30	19.80
2000	Annual	30.19	22.56
1999	Annual	29.87	15.81

On February 27, 2004, the closing sales price per Ordinary ADS on the NYSE was \$22.35.

The table below sets forth, for the periods indicated, the high and low closing sales prices for the Preference ADSs on the NYSE:

		Price per preference ADS (\$)	
		High	Low
2004	February	16.05	14.95
	January	17.20	14.89
2003	December	16.68	15.20
	November	15.90	13.85
	October	15.11	13.74
	September	15.00	12.40
	August	12.44	11.50
2003	Fourth Quarter	16.68	13.74
	Third Quarter	15.00	11.50
	Second Quarter	12.60	10.90
	First Quarter	12.35	9.85
2002	Fourth Quarter	11.10	5.30
	Third Quarter	12.11	4.90
	Second Quarter	15.70	10.90
	First Quarter	15.68	12.46
2003	Annual	16.68	9.85
2002	Annual	15.70	4.90

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2001	Annual	19.64	14.00
2000	Annual	16.91	13.25
1999	Annual	16.75	11.25

On February 27, 2004, the closing sales price per Preference ADS on the NYSE was \$15.15.

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Item 10. Additional information

B. Articles of Association

Fresenius Medical Care AG is a stock corporation (Aktiengesellschaft) organized under the laws of Germany. Fresenius Medical Care AG is registered with the commercial register of the local court (*Amtsgericht*) of Hof an der Saale, Germany under HRB 2460. Our registered office (*Sitz*) is Hof an der Saale, Germany. Our business address is Else-Kröner-Strasse 1, 61352 Bad Homburg, Germany, telephone +49-6172-609-0.

Corporate Purposes

Under our articles of association, our corporate purposes are:

developing, producing, distributing and selling, health care products, systems and procedures, primarily dialysis products and systems;

planning, establishing, acquiring and operating health care businesses, including, but not limited to, dialysis clinics, directly or through third parties and through participation in joint ventures and other entities;

developing, producing and distributing other pharmaceutical products and the provision of health care services;

providing advice in the medical and pharmaceutical areas and disseminating scientific information and documentation; and

providing laboratory services for dialysis and non-dialysis patients and home health services.

We conduct our business directly and through subsidiaries within and outside Germany.

Voting Rights

Each Ordinary share entitles the holder thereof to one vote at general meetings of our shareholders. Resolutions are passed at a general or special meeting of our shareholders by a majority of the votes cast, unless a higher vote is required by law or our articles of association.

Our Preference shares do not have any voting rights, except as described in this paragraph. If we do not pay the minimum annual dividend payable on the Preference shares for any year in the following year, and we do not pay both the dividend arrearage and the dividend payable on the Preference shares for the following year in full in the next following year, then the Preference shares shall have the same voting rights as the Ordinary shares (one vote for each share held or for each three ADSs held) until all Preference share dividend arrearages are fully paid up. In addition, holders of Preference shares are entitled to vote on any matters affecting their preferential rights, such as changes in the rate of the preferential dividend. Any such vote requires the affirmative vote of 75% of the votes cast in a meeting of holders of Preference shares.

Dividend Rights

Our management board and supervisory board will propose any dividends for approval at the annual general meeting of shareholders, which must be held within the first eight months following the end of a fiscal year. Usually the shareholders vote on a recommendation made by our management board and our supervisory board as to the amount of dividend to be paid. Any dividends are paid once a year.

Under German law, dividends are payable from the prior year unconsolidated retained earnings of Fresenius Medical Care AG, determined in accordance with German accounting principles (*Bilanzgewinn*).

Dividends are paid on presentation of the relevant dividend coupon to us or to our paying agent or agents appointed by us from time to time. If the Ordinary shares and the Preference shares which are entitled to dividend payments are held in a clearing system, the dividends will be paid in accordance with the rules of the individual clearing system. We will publish notice of the dividends paid and the appointment of the paying agent or agents for this purpose in the German Federal Gazette (*Bundesanzeiger*).

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In the case of holders of ADRs, the depositary will receive all dividends and distributions on all deposited securities and will, as promptly as practicable, distribute the dividends and distributions to the holders of ADRs entitled to the dividend. See Description of American Depositary Receipts Share Dividends and Other Distributions.

For each fiscal year, our management board and the supervisory board propose the treatment of all unappropriated profits, including the amount of our net profits which will be distributed by way of dividends, subject to shareholder approval. The management board and the supervisory board may allocate up to 50% of unappropriated profits to our free reserve (that is, they may determine not to distribute such profits) without shareholder approval, in which case the shareholders approve the treatment of the balance of such profits. Under German law, we must pay the annual dividend paid on the Preference shares, in all cases, before we pay any dividend on the Ordinary shares.

Liquidation Rights

In accordance with the German Stock Corporation Act (*Aktiengesetz*), upon a liquidation, any liquidation proceeds remaining after paying all of our liabilities will be distributed among the holders of Preference shares and the holders of Ordinary shares in proportion to the total number of the shares held by each holder. The Preference shares are not entitled to a preference in liquidation.

Preemptive Rights

Under the German Stock Corporation Act, an existing stockholder in a stock corporation, including a holder of Preference shares, has a preferential right to subscribe for any issue by that corporation of shares, debt instruments convertible into shares and participating debt instruments in proportion to the number of shares held by that stockholder in the existing capital of the corporation. The German Stock Corporation Act provides that this preferential right can only be excluded by a resolution of the general meeting passed at the same time as the resolution authorizing the capital increase. A supermajority of at least three quarters of the share capital represented at the general meeting is required for the exclusion. The waiver of the preemptive rights of the holders of Preference shares requires a vote of these holders of 75% of the capital represented by Preference shares at the meeting at which the vote is taken. In addition, a special justification by the corporation stating that the goal pursued by the corporation with the issuance of the new security could not reasonably be achieved without and outweighs the elimination of preemptive rights, is generally required for the exclusion. A special justification is not required for any increase in the share capital for contributions in cash if the increase does not exceed 10% of the existing capital and the issue price is not materially less than the price for the shares quoted on a stock exchange.

General Meeting

Our annual general meeting must be held within the first eight months of each fiscal year at the location of Fresenius Medical Care AG's registered office, or in a German city where a stock exchange is situated or at the location of a registered office of a domestic affiliated company. Each of our shareholders is entitled to participate in a general meeting regardless of whether they possess voting rights.

Description of American Depositary Receipts

Morgan Guaranty Trust Company of New York, a New York banking corporation, is the depositary for our Ordinary shares and our Preference shares. Each ADS represents an ownership interest in one-third of one Ordinary share or one Preference share. We deposit the underlying shares with the custodian, as agent of the depositary, under the deposit agreements among ourselves, the depositary and all of the ADS holders of the applicable class. Each ADS also represents any securities, cash or other property deposited with the depositary but not distributed by it directly to ADS holders. The ADSs are evidenced by securities called American depositary receipts or ADRs. An ADR may be issued in either book-entry or certificated form by the depositary. If an ADR is issued in book-entry form, owners will receive periodic statements from the depositary showing their ownership interest in ADSs.

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The depositary's office is located at 60 Wall Street, New York, NY 10260.

An investor may hold ADSs either directly or indirectly through a broker or other financial institution. Investors who hold ADSs directly, by having an ADS registered in their names on the books of the depositary, are ADR holders. This description assumes an investor holds ADSs directly. Investors who hold ADSs through their brokers or financial institution nominees must rely on the procedures of their brokers or financial institutions to assert the rights of an ADR holder described in this section. Investors should consult with their brokers or financial institutions to find out what those procedures are.

Because the depositary's nominee will actually be the registered owner of the shares, investors must rely on it to exercise the rights of a shareholder on their behalf. The obligations of the depositary and its agents are set out in the deposit agreement. The deposit agreement and the ADSs are governed by New York law.

The following is a summary of the material terms of the deposit agreements. Because it is a summary, it does not contain all the information that may be important to investors. Except as specifically noted, the description covers both Ordinary ADSs and Preference ADSs. For more complete information, investors should read the entire applicable deposit agreement and the form of ADR of the relevant class which contains the terms of the ADSs. Investors may obtain a copy of the deposit agreements at the SEC's Public Reference Room which is located at 450 Fifth Street, N.W., Washington, D.C. 20549.

Share Dividends and Other Distributions

We may make various types of distributions with respect to our Ordinary shares and our Preference shares. The depositary has agreed to pay to investors the cash dividends or other distributions it or the custodian receives on the shares or other deposited securities, after deducting its expenses. Investors will receive these distributions in proportion to the number of underlying shares of the applicable class their ADSs represent.

Except as stated below, to the extent the depositary is legally permitted it will deliver distributions to ADR holders in proportion to their interests in the following manner:

Cash. The depositary shall convert cash distributions from foreign currency to U.S. dollars if this is permissible and can be done on a reasonable basis. The depositary will endeavor to distribute cash in a practicable manner, and may deduct any taxes or other governmental charges required to be withheld, any expenses of converting foreign currency and transferring funds to the United States, and certain other expenses and adjustments. In addition, before making a distribution the depositary will deduct any taxes withheld. If exchange rates fluctuate during a time when the depositary cannot convert a foreign currency, investors may lose some or all of the value of the distribution.

Shares. If we make a distribution in shares, the depositary will issue additional ADRs to evidence the number of ADSs representing the distributed shares. Only whole ADSs will be issued. Any shares which would result in fractional ADSs will be sold and the net proceeds will be distributed to the ADR holders otherwise entitled to receive fractional ADSs.

Rights to receive additional shares. In the case of a distribution of rights to subscribe for Ordinary shares, Preference shares or other rights, if we provide satisfactory evidence that the depositary may lawfully distribute the rights, the depositary may arrange for ADR holders to instruct the depositary as to the exercise of the rights. However, if we do not furnish the required evidence or if the depositary determines it is not practical to distribute the rights, the depositary may

sell the rights if practicable and distribute the net proceeds as cash, or

allow the rights to lapse, in which case ADR holders will receive nothing.

We have no obligation to file a registration statement under the Securities Act in order to make any rights available to ADR holders.

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Other Distributions. If we make a distribution of securities or property other than those described above, the depositary may either:

distribute the securities or property in any manner it deems fair and equitable;

after consultation with us if practicable, sell the securities or property and distribute any net proceeds in the same way it distributes cash; or

hold the distributed property in which case the ADSs will also represent the distributed property.

Any dollars will be distributed by checks drawn on a bank in the United States for whole dollars and cents (fractional cents will be withheld without liability for interest and added to future cash distributions).

The depositary may choose any practical method of distribution for any specific ADR holder, including the distribution of foreign currency, securities or property, or it may retain the items, without paying interest on or investing them, on behalf of the ADR holder as deposited securities.

The depositary is not responsible if it decides that it is unlawful or impractical to make a distribution available to any ADR holders.

There can be no assurance that the depositary will be able to convert any currency at a specified exchange rate or sell any property, rights, shares or other securities at a specified price, or that any of these transactions can be completed within a specified time period.

Deposit, Withdrawal and Cancellation

The depositary will issue ADSs if an investor or his broker deposits Ordinary shares or Preference shares or evidence of rights to receive Ordinary shares or Preference shares with the custodian. Shares deposited with the custodian must be accompanied by certain documents, including instruments showing that such shares have been properly transferred or endorsed to the person on whose behalf the deposit is being made.

The custodian will hold all deposited shares for the account of the depositary. ADR holders thus have no direct ownership interest in the shares and only have the rights that are contained in the deposit agreement. The custodian will also hold any additional securities, property and cash received on or in substitution for the deposited shares. The deposited shares and any additional items are referred to as deposited securities.

Upon each deposit of shares, receipt of related delivery documentation and compliance with the other provisions of the deposit agreement, including the payment of the fees and charges of the depositary and any taxes or other fees or charges owing, the depositary will issue an ADR or ADRs of the applicable class in the name of the person entitled to them. The ADR or ADRs will evidence the number of ADSs to which the person making the deposit is entitled. Certificated ADRs will be delivered at the depositary's principal New York office or any other location that it may designate as its transfer office.

All ADSs issued will, unless specifically requested to the contrary, be part of the depositary's book-entry direct registration system, and a registered holder will receive periodic statements from the depositary which will show the number of ADSs registered in the holder's name. An ADR holder can request that the ADSs not be held through the depositary's direct registration system and that a certificated ADR be issued. If ADRs are in book-entry form, a statement setting forth the holder's ownership interest will be mailed to holders by the depositary.

When an investor surrenders ADSs at the depositary's office, the depositary will, upon payment of certain applicable fees, charges and taxes, and upon receipt of proper instructions, deliver the whole number of shares of the applicable class represented by the ADSs turned in to the account the investor directs within Clearstream Banking AG, the central German clearing firm.

The depositary may only restrict the withdrawal of deposited securities in connection with:

temporary delays caused by closing our transfer books or those of the depositary, or the deposit of shares in connection with voting at a shareholders' meeting, or the payment of dividends,

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the payment of fees, taxes and similar charges, or

compliance with any U.S. or foreign laws or governmental regulations relating to the ADRs.

This right of withdrawal may not be limited by any other provision of the applicable deposit agreement.

Voting Rights

Only the depositary's nominee is able to exercise voting rights with respect to deposited shares. Upon receipt of a request from the depositary for voting instructions, a holder of ADSs may instruct the depositary how to exercise the voting rights for the shares which underlie their ADSs. After receiving voting materials from us, the depositary will notify the ADR holders of any shareholder meeting or solicitation of consents or proxies. This notice will describe how holders may instruct the depositary to exercise voting rights for the shares which underlie their ADSs. For instructions to be valid, the depositary must receive them on or before the date specified in the depositary's request for instructions. The depositary will try, as far as is practical, subject to the provisions of and governing the underlying shares or other deposited securities, to vote or to have its agents vote the shares or other deposited securities as instructed. The depositary will only vote or attempt to vote as holders instruct. The depositary will not itself exercise any voting discretion. Neither the depositary nor its agents are responsible for any failure to carry out any voting instructions, for the manner in which any vote is cast or for the effect of any vote.

Our Preference shares are non-voting, except in a limited number of circumstances. In those circumstances in which Preference shares are entitled to vote, the procedures and limitations described above will apply in connection with the depositary's request for voting instructions from holders of ADSs representing Preference shares.

There is no guarantee that holders will receive voting materials in time to instruct the depositary to vote and it is possible that holders, or persons who hold their ADSs through brokers, dealers or other third parties, will not have the opportunity to exercise a right to vote.

Description of the Pooling Agreements

The information under the heading "DESCRIPTION OF THE POOLING AGREEMENTS" set forth in the prospectus of Fresenius Medical Care AG dated July 20, 2000 is incorporated herein by reference.

C. Material contracts

For information regarding certain of our material contracts, see "Item 7.B. Major Shareholders and Related Party Transactions" "Related Party Transactions." For a description of our stock option plans, see "Item 6.E. Directors, Senior Management and Employees" "Share Ownership Options to Purchase our Securities." For a description of our 2003 Senior Credit Agreement, see "Item 5B. Operating and Financial Review and Prospects" "Liquidity and Capital Resources." Our material agreements also include the agreements that FMCH and certain of its subsidiaries entered into with the U.S. government when we settled a U.S. government investigation. Our Report on Form 6-K filed with the SEC on January 27, 2000 contains a description of the agreements comprising the settlement, including the plea agreements and a corporate integrity agreement in Part II, Item 5 "Other Events" "OIG Investigation," which is incorporated herein by reference.

Our material agreements include the settlement agreement that we, FMCH and NMC entered into with the Official Committee of Asbestos Injury Claimants, and the Official Committee of Asbestos Property Damage Claimants of W.R. Grace & Co. A description of the terms of the settlement agreement appears in Item 8.A.7 "Legal Proceedings."

D. Exchange controls

Exchange Controls and Other Limitations Affecting Security Holders.

At the present time, Germany does not restrict the export or import of capital, except for investments in areas like Iraq, Serbia, Montenegro or Sierra Leone. However, for statistical purposes only, every resident individual or

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corporation residing in Germany must report to the German Federal Bank (*Deutsche Bundesbank*), subject only to certain immaterial exceptions, any payment received from or made to an individual or a corporation resident outside of Germany if such payment exceeds 12,500. In addition, residents must report any claims against, or any liabilities payable to, non-residents individuals or corporations, if such claims or liabilities, in the aggregate 5 million at the end of any month.

There are no limitations imposed by German law or our articles of association (*Satzung*) on the right of a non-resident to hold the Preference shares or the ADSs evidencing Preference shares or Ordinary shares.

E. Taxation

The following is a discussion of the material United States federal income and German tax consequences to Qualified Holders holding Fresenius Medical Care Ordinary shares, preference shares or ADSs with respect to such shares (collectively the shares). This discussion is based upon existing United States federal income and German tax law, including legislation, regulations, administrative rulings and court decisions, as in effect on the date of this Annual Report, all of which are subject to change, possibly with retroactive effect. For purposes of this discussion, in general, a Qualified Holder means a beneficial owner of Fresenius Medical Care shares that (1) is a resident of the United States for purposes of the United States-Germany income tax treaty (the Income Tax Treaty), which generally includes an individual United States resident, a corporation created or organized under the laws of the United States, any state thereof or the District of Columbia and a partnership, estate or trust, to the extent its income is subject to taxation in the United States as the income of a United States resident, either in its hands or in the hands of its partners or beneficiaries, (2) does not hold Fresenius Medical Care shares as part of the business property of a permanent establishment located in Germany or as part of a fixed base of an individual located in Germany and used for the performance of independent personal services and (3) if not an individual, is not subject to the limitation on benefits restrictions in the Income Tax Treaty. This discussion assumes that the Qualified Holder holds Fresenius Medical Care shares as a capital asset. This discussion does not address all aspects of United States federal income and German taxation that may be relevant to all Qualified Holders in light of their particular circumstances, including for example Qualified Holders whose stock was acquired pursuant to the exercise of an employee stock option or otherwise as compensation or Qualified Holders who are subject to special treatment under United States federal income tax laws (for example, financial institutions, insurance companies, tax-exempt organizations and broker-dealers). This discussion also does not address any aspects of state, local or non-United States (other than certain German) tax law.

EACH QUALIFIED HOLDER IS STRONGLY URGED TO CONSULT HIS OR HER TAX ADVISOR AS TO THE UNITED STATES FEDERAL INCOME AND GERMAN TAX CONSEQUENCES OF HOLDING FRESENIUS MEDICAL CARE SHARES, INCLUDING THE PARTICULAR FACTS AND CIRCUMSTANCES THAT MAY BE UNIQUE TO SUCH QUALIFIED HOLDER, AND AS TO ANY OTHER TAX CONSEQUENCES OF HOLDING OUR SHARES.

Taxation of Dividends

For dividends distributed in 2004 out of profits earned in or before 2003, German corporations are required to withhold German tax on dividends in an amount equal to 20% of the gross amount paid to resident and non-resident stockholders. A partial refund of this 20% withholding tax can be obtained by Qualified Holders under the Income Tax Treaty (subject to certain limitations). Qualified Holders are generally subject to United States federal income tax on dividends paid by German corporations. Subject to applicable limitations of United States federal income tax law, Qualified Holders may be able to claim a foreign tax credit for certain German income taxes paid. The amount of the refund of German withholding tax and the determination of the foreign tax credit allowable against United States federal income tax generally depend on whether or not the Qualified Holder is a United States corporation owning at least 10% of the voting stock of Fresenius Medical Care AG (a 10% Holder).

In the case of any Qualified Holder other than a 10% Holder, the German withholding tax on the dividends paid in 2004 is partially refunded under the Income Tax Treaty, effectively reducing the withholding tax to 15% of the gross amount of the dividend. Thus, for each \$100 of gross dividend paid by Fresenius Medical Care AG in

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2004 to a Qualified Holder (other than a 10% Holder), the dividend after partial refund of the 20% withholding tax under the Income Tax Treaty will be subject to a German withholding tax of \$15.

In the case of a 10% Holder, the 20% German withholding tax on dividends paid in 2004 is reduced under the Income Tax Treaty to 5% of the gross amount of the dividend. Such 10% Holders may, therefore, apply for a refund of German withholding tax on the dividend paid in 2002 in the amount of 15% of the gross amount of the dividend. Subject to applicable limitations of United States federal income tax laws, a 10% Holder may be entitled to a foreign tax credit for the 5% German withholding tax on dividends and for the portion of the total income taxes (trade income tax and corporation income tax, including any surtax) paid by Fresenius Medical Care AG attributable to distributed profits.

The German corporate imputation system that has provided German resident individual shareholders with a tax credit in respect of dividends paid by German corporations was recently repealed, effective with respect to dividends paid after 2002. Consequently, Qualified Holders are no longer be entitled to the additional 5% treaty refund with respect to such dividends. However, the German withholding tax was reduced to 20% with respect to such dividends.

Dividends paid in euros to a Qualified Holder of Fresenius Medical Care shares will be included in income in a dollar amount calculated by reference to the exchange rate in effect on the date the dividends (including any deemed refund of German corporate tax) are received or treated as received by such holder. If dividends paid in euros are converted into dollars on the date received or treated as received, Qualified Holders generally should not be required to recognize foreign currency gain or loss in respect of each dividend.

A surtax on the German withholding tax is currently levied on dividend distributions paid by a German resident company. The rate of this surtax is 5.5%, which is 1.110% (5.5% 2/3 20%) of the gross dividend amount. Under the Income Tax Treaty, Qualified Holders are entitled to a full refund of this surtax.

Refund Procedures

To claim the refund reflecting the current reduction of the German withholding tax from 20% to 15%, the additional 5% treaty refund and the refund of the effective 1.110% German surtax, when applicable, a Qualified Holder must submit (either directly or, as described below, through the U.S. transfer agent for Fresenius Medical Care shares or the Depository Trust Company) a claim for refund to the German tax authorities, with the original bank voucher (or certified copy thereof) issued by the paying entity documenting the tax withheld within four years from the end of the calendar year in which the dividend is received. Claims for refunds are made on a special German claim for refund form, which must be filed with the German tax authorities: Bundesamt für Finanzen, 53221 Bonn-Beuel, Germany. The German claim for refund forms may be obtained from the German tax authorities at the same address where the applications are filed or from the Embassy of the Federal Republic of Germany, 4645 Reservoir Road, N.W., Washington, D.C. 20007-1998.

Qualified Holders must also submit to the German tax authorities certification (IRS Form 6166) of their last filed United States federal income tax return. Such certification is obtained from the office of the Director of the Internal Revenue Service Center by filing a request for the certification with the Internal Revenue Service Philadelphia Service Center, Foreign Certification Request, P.O. Box 16347, Philadelphia, PA 19114-0447. (Additional information, including IRS Publication 686, can be obtained from the Internal Revenue Service website at www.irs.gov.) Requests for certification are to be made in writing and must include the Qualified Holder's name, social security number or employer identification number, tax return form number and tax period for which certification is requested. The Internal Revenue Service will send the certification directly to the German tax authorities if the Qualified Holder authorizes the Internal Revenue Service to do so. This certification is valid for three years and need only be resubmitted in a fourth year in the event of a subsequent application for refund.

The U.S. transfer agent will receive and distribute dividends to Qualified Holders who hold Fresenius Medical Care shares of record and will perform administrative functions necessary to claim the refund reflecting the current reduction in German withholding tax from 20% to 15% (to 5% for 10% Holders), the additional 5%

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treaty refund and the refund of the effective 1.110% German surtax, when applicable, for such holders. These arrangements may be amended or revoked at any time in the future.

Under a newly implemented simplified and accelerated refund procedure, the U.S. transfer agent will prepare the German claim for refund forms on behalf of Qualified Holders and file them electronically with the German tax authorities. In order for the U.S. transfer agent to file the claim for refund forms, the U.S. transfer agent will prepare and mail to these Qualified Holders, and the holders will be requested to sign and return to the U.S. transfer agent, (1) a statement authorizing the U.S. transfer agent to perform these procedures and agreeing that the German tax authorities may inform the IRS of any refunds of German taxes and (2) a written authorization to remit the refund of withholding to an account other than that of the Qualified Holder. The U.S. transfer agent will attach the signed statement and the documentation issued by the paying agency documenting the dividend paid and the tax withheld to the claim for refund form and file them with the German tax authorities. Qualified Holders should also request certification (IRS Form 6166) of their last filed United States federal income tax return from the IRS and have it ready for presentation to the U.S. transfer agent upon request. If the Qualified Holder fails to present this certification within a reasonable time after the request, the refund of the German withholding taxes will be denied.

A simplified refund procedure for Qualified Holders whose Fresenius Medical Care shares are registered with brokers participating in the Depository Trust Company is in effect between the Depository Trust Company and the German tax authorities. Under this simplified refund procedure, the Depository Trust Company provides the German tax authorities with electronic certification of the U.S. taxpayer status of such Qualified Holders based on information it receives from its broker participants, and claims a refund on behalf of those Qualified Holders. Accordingly, these Qualified Holders do not need to file refund claim forms through the U.S. transfer agent.

The German tax authorities will issue refunds denominated in euros. The refunds will be issued in the name of the U.S. transfer agent or the Depository Trust Company, as the case may be, which will convert the refunds to dollars and make corresponding refund payments to Qualified Holders and to brokers. The brokers, in turn, will remit corresponding refund amounts to the Qualified Holders holding Fresenius Medical Care shares registered with such brokers. Qualified Holders of Fresenius Medical Care shares who receive a refund attributable to reduced withholding taxes under the Income Tax Treaty may be required to recognize foreign currency gain or loss, which will be treated as income or loss, to the extent that the dollar value of the refund received or treated as received by the Qualified Holder differs from the U.S. dollar equivalent of the refund on the date the dividend on which such withholding taxes were imposed was received or treated as received by the Qualified Holder.

Taxation of Capital Gains

Under the Income Tax Treaty, a Qualified Holder will not be liable for German tax on capital gains realized or accrued on the sale or other disposition of Fresenius Medical Care shares.

Upon a sale or other disposition of Fresenius Medical Care shares, a Qualified Holder will recognize capital gain or loss for United States federal income tax purposes equal to the difference between the amount realized and the Qualified Holder's adjusted tax basis in the Fresenius Medical Care shares. In the case of an individual Qualified Holder of Fresenius Medical Care shares, any such capital gain will be subject to a maximum United States federal income tax rate of 15%, if the individual Qualified Holder's holding period in the Fresenius Medical Care shares is more than 12 months.

German Gift and Inheritance Taxes

Under the estate, inheritance and gift tax treaty between the United States and Germany (the Estate Tax Treaty), a transfer of shares or ADS's generally will not be subject to the German gift or inheritance tax so long as neither the donor or decedent, nor heir, donee or other beneficiary, was domiciled in Germany for the purpose of the Estate Tax Treaty at the time of the transfer, and the shares or ADSs were not held as part of a permanent base of fixed establishment in Germany

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The United States-Germany estate tax treaty also provides a credit against United States federal estate and gift tax liability for the amount of inheritance and gift tax paid in Germany, subject to certain limitations, in a case where the Fresenius Medical Care shares are subject to German inheritance or gift tax and United States federal estate or gift tax.

United States Information Reporting and Backup Withholding

Dividends on Fresenius Medical Care shares, and payments of the proceeds of a sale of Fresenius Medical Care shares, paid within the United States or through certain U.S.-related financial intermediaries are subject to information reporting and may be subject to backup withholding at a 28% rate unless the Qualified Holder (1) is a corporation or other exempt recipient or (2) provides a taxpayer identification number and certifies that no loss of exemption from backup withholding has occurred.

H. Documents on display

We file periodic reports and information with the Securities and Exchange Commission and the New York Stock Exchange. You may inspect a copy of these reports without charge at the Public Reference Room of the Securities and Exchange Commission at Room 1024, 450 Fifth Avenue, N.W., Washington, D.C. 20549 or at the Securities and Exchange Commission's regional offices 233 Broadway, New York, New York 10279 and 500 West Madison Street, Suite 1400, Chicago, Illinois 60661. The public may obtain information on the operation of the Public Reference Room by calling the Securities and Exchange Commission at 1-800-SEC-0330. The Securities and Exchange Commission also maintains an Internet site that contains reports, proxy and information statements and other information regarding registrants that file electronically with the Securities and Exchange Commission. The Securities and Exchange Commission's World Wide Web address is <http://www.sec.gov>.

The New York Stock Exchange currently lists American Depositary Shares representing our Preference shares and American Depositary Shares representing our Ordinary shares. As a result, we are subject to the periodic reporting requirements of the Securities Exchange Act of 1934, as amended, and we file reports and other information with the Securities and Exchange Commission. These reports, proxy statements and other information and the registration statement and exhibits and schedules thereto may be inspected without charge at, and copies thereof may be obtained at prescribed rates from, the public reference facilities of the Securities and Exchange Commission and the electronic sources listed in the preceding paragraph. In addition, these materials are available for inspection and copying at the offices of the New York Stock Exchange, 20 Broad Street, New York, New York 1005, USA.

We prepare annual and quarterly reports, which are then distributed to our shareholders. Our annual reports contain financial statements examined and reported upon, with opinions expressed by our independent auditors. The consolidated financial statements of Fresenius Medical Care included in these annual reports are prepared in conformity with U.S. generally accepted accounting principles. Our annual and quarterly reports to our shareholders are posted on our website at <http://www.fmc-ag.com>. In furnishing our web site address in this report, however, we do not intend to incorporate any information on our web site with this report, and any information on our web site should not be considered to be part of this report.

We will also furnish the depositary with all notices of shareholder meetings and other reports and communications that are made generally available to our shareholders. The depositary, to the extent permitted by law, shall arrange for the transmittal to the registered holders of American Depositary Receipts of all notices, reports and communications, together with the governing instruments affecting the Preference shares and any amendments thereto, available for inspection by registered holders of American Depositary Receipts at the principal office of the depositary, presently located at 60 Wall Street, New York, New York, 10260, USA.

Documents referred to in this report which related to us as well as future annual and interim reports prepared by us may also be inspected at our offices, Else-Kröner-Strasse 1, 61352 Bad Homburg.

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Item 11. *Quantitative and Qualitative Disclosures About Market Risk*

Market Risk

Our businesses operate in highly competitive markets and are subject to changes in business, economic and competitive conditions. Our business is subject to:

changes in reimbursement rates;

intense competition;

foreign exchange rate fluctuations;

varying degrees of acceptance of new product introductions;

technological developments in our industry;

uncertainties in litigation or investigative proceedings and regulatory developments in the health care sector; and

the availability of financing.

Our business is also subject to other risks and uncertainties that we describe from time to time in our public filings. Developments in any of these areas could cause our results to differ materially from the results that we or others have projected or may project.

Reimbursement Rates

We obtained approximately 43% of our worldwide revenue for 2003 from sources subject to regulations under U.S. government health care programs. In the past, U.S. budget deficit reduction and health care reform measures have changed the reimbursement rates under these programs, including the Medicare composite rate, the reimbursement rate for EPO, and the reimbursement rates for other dialysis and non-dialysis related services and products, as well as other material aspects of these programs, and they may change in the future.

We also obtain a significant portion of our net revenues from reimbursement by non-government payors. Historically, these payors reimbursement rates generally have been higher than government program rates in their respective countries. However, non-governmental payors are imposing cost containment measures that are creating significant downward pressure on reimbursement levels that we receive for our services and products.

Inflation

The effects of inflation during the periods covered by the consolidated financial statements have not been significant to our results of operations. However, most of our net revenues from dialysis care are subject to reimbursement rates regulated by governmental authorities, and a significant portion of other revenues, especially revenues from the U.S., is received from customers whose revenues are subject to these regulated reimbursement rates. Non-governmental payors are also exerting downward pressure on reimbursement rates. Increased operation costs that are subject to inflation, such as labor and supply costs, may not be recoverable through price increases in the absence of a compensating increase in reimbursement rates payable to us and our customers, and could materially adversely affect our business, financial condition and results of operations.

Management of Currency and Interest Rate Risks

We are primarily exposed to market risk from changes in foreign currency exchange rates and changes in interest rates. In order to manage the risks from these foreign currency exchange rate and interest rate fluctuations, we enter into various hedging transactions with investment grade financial institutions as authorized by the management board. We do not contract for financial instruments for trading or other speculative purposes.

We conduct our financial instrument activity under the control of a single centralized department. We have established guidelines for risk assessment procedures and controls for the use of financial instruments. They

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include a clear segregation of duties with regard to execution on one side and administration, accounting and controlling on the other.

Foreign Currency Exposure

We conduct our business on a global basis in several major international currencies, although our operations are primarily in the United States and Germany. For financial reporting purposes, we have chosen the U.S. dollar as our reporting currency. Therefore, changes in the rate of exchange between the U.S. dollar, the euro and the local currencies in which the financial statements of our international operations are maintained, affect our results of operations and financial position as reported in our consolidated financial statements. We have translated the balance sheets of our non-U.S. dollar denominated operations into U.S. dollars at the exchange rates prevailing at the balance sheet date. Revenues and expenses are translated at the average exchange rates for the period.

Our exposure to market risk for changes in foreign exchange rates relates to transactions such as sales and purchases, and lendings and borrowings, including intercompany borrowings. We sell significant amounts of products from our manufacturing facilities in Germany to our other international operations. In general, our German sales are denominated in euro. Consequently, our subsidiaries are exposed to fluctuations in the rate of exchange between the euro and the currency in which their local operations are conducted. We employ, to a limited extent, forward contracts and options to hedge our currency exposures. Our policy, which has been consistently followed, is that forward currency contracts and options be used only for hedging foreign currency exposures.

Our foreign exchange contracts contain credit risk, in that our bank counterparties may be unable to meet the terms of the agreements. We monitor the potential risk of loss with any one party from this type of risk. Our management does not expect any material losses as a result of default by the other parties. The table below provides information about our foreign exchange forward contracts at December 31, 2003. The information is provided in U.S. dollar equivalent amounts. The table presents the notional amounts by year of maturity, the fair values of the contracts, which show the unrealized net gain (loss) on existing contracts as of December 31, 2003, and the credit risk inherent to those contracts with positive market values as of December 31, 2003. All contracts expire within 36 months after the reporting date.

Foreign Currency Risk Management**December 31, 2003****(USD in thousands)**

	Nominal amount			total	Fair value	Credit risk
	2004	2005	2006			
Purchase of EUR against USD	502,273	14,427	359,743	876,443	91,858	91,858
Sale of EUR against USD	5,908			5,908	(428)	
Purchase of EUR against others	240,798	21,685		262,483	10,759	11,738
Sale of EUR against others	29,529			29,529	66	234
Others	34,331	8,661		42,992	(71)	951
Total	812,839	44,773	359,743	1,217,355	102,184	104,781

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A summary of the high and low exchange rates for the Euro to U.S. dollars and the average exchange rates for the last five years is set forth below. The Deutsche Mark (DM) was replaced by the euro () in the foreign exchange markets beginning in 1999 at a fixed conversion rate of DM 1.95583 = 1.

Year ending December 31,	Year s High	Year s Low	Year s Average	Year s Close
1999 \$ per	1.1790	1.0015	1.0658	1.0046
2000 \$ per	1.0388	0.8252	0.9236	0.9305
2001 \$ per	0.9545	0.8384	0.8956	0.8813
2002 \$ per	1.0487	0.8578	0.9454	1.0487
2003 \$ per	1.2630	1.0377	1.1312	1.2630

Interest Rate Exposure

We are exposed to changes in interest rates that affect our variable-rate based borrowings. We enter into debt obligations and into accounts receivable financings to support our general corporate purposes including capital expenditures and working capital needs. We enter into derivatives, particularly interest rate swaps, to protect interest rate exposures arising from long-term and short-term borrowings and our accounts receivable securitization programs at floating rates by effectively swapping them into fixed rates. Under interest rate swaps, we agree with other parties to exchange, at specified intervals, the difference between fixed-rate and floating-rate interest amounts calculated by reference to an agreed notional amount.

Our subsidiary, National Medical Care, has entered into dollar interest rate swap agreements with various commercial banks for notional amounts totaling \$950 million as of December 31, 2003. National Medical Care entered into all of these agreements for purposes other than trading.

The dollar interest rate swaps effectively change National Medical Care's interest rate exposure on the majority of its variable-rate loans under our senior credit agreement (\$912 million outstanding as of December 31, 2003), loans extended to us by Fresenius AG (\$30 million outstanding as of December 31, 2003), and the drawdowns under our receivables financing facility (drawn as of December 31, 2003, \$158 million) to a fixed interest rate of 5.45%. Our accounts receivable financing facility has been reflected in our consolidated financial statements as a reduction to accounts receivable.

The dollar interest rate swap agreements expire at various dates between January 2004 and December 2009. At December 31, 2003, the fair value of these agreements is \$(71.3) million.

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The table below presents principal amounts and related weighted average interest rates by year of maturity for the various dollar interest rate swap agreements and for our significant fixed-rate long-term debt obligations.

Dollar Interest Rate Exposure

December 31, 2003

(U.S. dollars in millions)

	2004	2005	2006	2007	2008	Thereafter	Totals	Fair Value Dec. 31, 2003
Principal payments on Senior Credit Agreement	68	104	104	254	4	378	\$ 912	\$ 912
Variable interest rate = 3.15%								
Interest rate swap agreements								
Notional amount	150		250	200	100	250	950	(71)
Average fixed pay rate = 5.45%	6.51%		4.60%	6.61%	4.86%	4.99%	5.45%	
Receive rate = 3-month \$LIBOR								
Company obligated mandatorily redeemable preferred securities of subsidiaries								
Fresenius Medical Care Capital Trusts								
Fixed interest rate = 7.875%/issued in 1998					450		450	477
Fixed interest rate = 7.375%/issued in 1998 (denominated in DEM)					194		194	205
Fixed interest rate = 7.875%/issued in 2001						225	225	241
Fixed interest rate = 7.375%/issued in 2001 (denominated in EUR)						379	379	402

Our subsidiaries FMC Japan and Fresenius Kawasumi have entered into Yen denominated interest rate swap agreements and a Yen-denominated interest rate cap agreement with a commercial bank for a notional amount of Yen 1,885 million as of December 31, 2003. The swaps change FMC Japan's and Fresenius Kawasumi's interest rate exposures on their variable-rate bank loans (Yen 1,249 million outstanding as of December 31, 2003) to a fixed interest rate of 2.99% on average. The Yen denominated interest rate swap agreements expire between March 2009 and June 2011. The cap agreement limits the interest rate risk for a notional amount of Yen 636 million as of December 31, 2003 to 2.8%. At December 31, 2003, the fair value of these agreements is \$(0.47) million. The terms of the Yen-denominated interest rate hedge agreements, especially the notional amounts outstanding at any specific point of time, match the terms of the bank loans which have been borrowed from the same bank that is counterparty in the swap and cap agreements. The amount of the bank borrowings and the notional amounts of both the swap agreements and the cap agreement always coincide until the final maturities when the bank debts are completely repaid and the swap and cap agreements expire.

Item 12. Description of Securities other than Equity Securities

Not applicable

PART II**Item 13. Defaults, Dividend Arrearages and Delinquencies**

None

Item 14. *Material Modifications to the Rights of Security Holders and Use of Proceeds*

None

Table of Contents**Item 15. Controls and Procedures**

The Company's management, including the Chief Executive Officer and Chief Financial Officer, have conducted an evaluation of the effectiveness of the Company's disclosure controls and procedures as of the end of the period covered by this report, as contemplated by Securities Exchange Act Rule 13a-15. Based on that evaluation, the Chief Executive Officer and Chief Financial Officer concluded that the disclosure controls and procedures are effective in ensuring that all material information required to be filed in this annual report has been made known to them in a timely fashion. There have been no significant changes in internal controls, or in factors that could significantly affect internal controls, subsequent to the date the Chief Executive Officer and Chief Financial Officer completed their evaluation.

Item 16A. Audit Committee Financial Expert

Although our supervisory board has not officially named a audit committee financial expert, three of the audit committee members could qualify under Item 16A of Form 20-F. All three of these audit committee members are independent as defined in Rule 10A-3 under the Securities Exchange Act of 1934, as amended.

Item 16B. Code of Ethics

In 2003 our management board adopted through our worldwide compliance program a code of ethics, titled the *Code of Business Conduct*, which applies to members of the management board, including its chairman and the responsible member for Finance & Controlling, other senior officers and all Company employees. It is filed with this report as an exhibit.

Item 16C. Principal Accountant Fees and Services

In the annual meeting held on May 22, 2003, our shareholders appointed KPMG Deutsche Treuhand-Gesellschaft AG Wirtschaftsprüfungsgesellschaft (KPMG), Berlin and Frankfurt am Main, to serve as our independent auditors for the 2003 fiscal year. KPMG billed the following fees to us for professional services in each of the last two fiscal years:

	2003	2002
	(in thousands)	
Audit fees	\$ 3,114	\$ 2,387
Audit related fees	329	194
Tax fees	834	1,237
Other fees	224	50
	—	—
Total	\$ 4,501	\$ 3,868

Audit Fees are the aggregate fees billed by KPMG for the audit of our consolidated and annual financial statements, reviews of interim financial statements and attestation services that are provided in connection with statutory and regulatory filings or engagements. **Audit-Related Fees** are fees charged by KPMG for assurance and related services that are reasonably related to the performance of the audit or review of our financial statements and are not reported under **Audit Fees**. This category comprises fees billed for the audit of employee benefit plans and pension schemes, agreed-upon procedure engagements and other attestation services subject to regulatory requirements, certifications of accounting-related internal controls, as well as advisory services associated with our financial reporting. **Tax Fees** are fees for professional services rendered by KPMG for tax compliance, tax advice on actual or contemplated transactions, tax consulting associated with international transfer prices, and expatriate employee tax services. Fees disclosed under the category **All Other Fees** are mainly related to our project **Internal Control over Financial Reporting** implementing the requirements of Section 404 of Sarbanes-Oxley Act of 2002. This category also includes training and other immaterial support services.

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Audit Committee s pre-approval policies and procedures

Our Audit Committee nominates and engages our independent auditors to audit our financial statements. See also the description in Item 6C. Directors, Senior management and Employees- Board Practices. In 2003 our Audit Committee also adopted a policy requiring management to obtain the Committee s approval before engaging our independent auditors to provide any audit or permitted non-audit services to us or our subsidiaries. Pursuant to this policy, which is designed to assure that such engagements do not impair the independence of our auditors, the Audit Committee pre-approves annually a catalog of specific audit and non-audit services in the categories Audit Services, Audit-Related Services, Tax Consulting Services, and Other Services that may be performed by our auditors as well as additional approval requirements based on fee amount.

Our Chief Financial Officer reviews all individual management requests to engage our auditors as a service provider in accordance with this catalog and, if the requested services are permitted pursuant to the catalog and fee level, approves the request accordingly. We inform the Audit Committee about these approvals on a quarterly basis. Services that are not included in the catalog or exceed applicable fee level require pre-approval by the Audit Committee s chairman or full Audit Committee on a case-by-case basis. Neither the chairman of our Audit Committee nor full Audit Committee is permitted to approve any engagement of our auditors if the services to be performed either fall into a category of services that are not permitted by applicable law or the services would be inconsistent with maintaining the auditors independence.

Item 16D. Exemptions from the Listing Standards for Audit Committees

Not applicable

Item 16E. Purchase of Equity Securities by the Issuer and Affiliated Purchasers

Not Applicable

PART III

Item 17. Financial Statements

Not applicable. See Item 18. Financial Statements.

Item 18. Financial Statements

The information called for by this item commences on Page F-1.

Item 19. Exhibits

Pursuant to the provisions of the Instructions for the filings of Exhibits to Annual Reports on Form 20-F, the Registrant is filing the following exhibits:

1.1 Amended Memorandum and Articles of Association (*Satzung*) of Fresenius Medical Care AG (Incorporated by reference to Exhibit 1.1 to the Registrant s Report on Form 6-K filed August 14, 2003).

2.1 Deposit Agreement between Morgan Guaranty Trust Company and Fresenius Medical Care AG dated August 5, 1996 relating to Ordinary Share ADSs (Incorporated by reference to Exhibit A to the Registrant s Registration Statement on Form F-6, Registration No. 333-5356, filed August 5, 1996).

2.2 Deposit Agreement between Morgan Guaranty Trust Company and Fresenius Medical Care AG dated as of November 22, 1996 relating to Preference Share ADSs (Incorporated by reference to Exhibit A to the Registrant s Registration Statement on Form F-6, Registration No. 333-5928, filed October 30, 1996).

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2.3 FMC Ordinary Shares Pooling Agreement dated September 27, 1996 by and between Fresenius AG, Fresenius Medical Care AG and the individuals acting from time to time as Independent Directors. (Incorporated by reference to Exhibit 10.1 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 4, 1996).

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2.4 FMC Preference Shares Pooling Agreement dated November 27, 1996, by and between Fresenius AG, Fresenius Medical Care AG, and the individuals acting from time to time as Independent Directors. (Incorporated by reference to Exhibit 2.7 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1996, filed April 7, 1997).

2.5 Senior Subordinated Indenture (U.S. Dollar denominated) dated as of February 19, 1998, among Fresenius Medical Care AG, FMC Trust Finance S.à.r.l. Luxembourg, State Street Bank and Trust Company, as Trustee, and the Subsidiary Guarantors named therein. (Incorporated by reference to Exhibit 2.6 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.6 Senior Subordinated Indenture (DM denominated) dated as of February 19, 1998, among Fresenius Medical Care AG, FMC Trust Finance S.à.r.l. Luxembourg, State Street Bank and Trust Company, as Trustee, and the Subsidiary Guarantors named therein. (Incorporated by reference to Exhibit 2.7 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.7 Declaration of Trust Establishing Fresenius Medical Care Capital Trust II, dated February 12, 1998. (Incorporated by reference to Exhibit 2.1 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.8 Declaration of Trust Establishing Fresenius Medical Care Capital Trust III, dated February 12, 1998. (Incorporated by reference to Exhibit 2.2 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.9 First Amendment to Declaration of Trust Establishing Fresenius Medical Care Capital Trust III, dated February 12, 1998. (Incorporated by reference to Exhibit 2.3 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.10 Amended and Restated Declaration of Trust of Fresenius Medical Care Capital Trust II, dated as of February 19, 1998. (Incorporated by reference to Exhibit 2.4 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.11 Amended and Restated Declaration of Trust of Fresenius Medical Care Capital Trust III, dated as of February 19, 1998. (Incorporated by reference to Exhibit 2.5 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.12 Guarantee Agreement dated as of February 19, 1998 between Fresenius Medical Care AG and State Street Bank and Trust Company as Trustee, with respect to Fresenius Medical Care Capital Trust II. (Incorporated by reference to Exhibit 2.8 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.13 Guarantee Agreement dated as of February 19, 1998 between Fresenius Medical Care AG and State Street Bank and Trust Company as Trustee, with respect to Fresenius Medical Care Capital Trust III. (Incorporated by reference to Exhibit 2.9 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.14 Agreement as to Expenses and Liabilities between Fresenius Medical Care AG and Fresenius Medical Care Capital Trust II dated as of February 19, 1998. (Incorporated by reference to Exhibit 2.10 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.15 Agreement as to Expenses and Liabilities between Fresenius Medical Care AG and Fresenius Medical Care Capital Trust III dated as of February 19, 1998. (Incorporated by reference to Exhibit 2.11 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1997, filed March 27, 1998).

2.16 Declaration of Trust of Fresenius Medical Care Capital Trust IV, dated February 12, 1998 (Incorporated by reference to Exhibit no. 4.41 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

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2.17 First Amendment to Declaration of Trust of Fresenius Medical Care Capital Trust IV, dated June 5, 2001 (Incorporated by reference to Exhibit No. 4.42 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.18 Declaration of Trust of Fresenius Medical Care Capital Trust V, dated June 1, 2001 (Incorporated by reference to Exhibit No. 4.43 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.19 Amended and Restated Declaration of Trust of Fresenius Medical Care Capital Trust IV, dated as of June 6, 2001 (Incorporated by reference to Exhibit No. 4.44 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.20 Amended and Restated Declaration of Trust of Fresenius Medical Care Capital Trust V, dated as of June 15, 2000 (Incorporated by reference to Exhibit No. 4.45 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.21 Senior Subordinated Indenture (U.S. Dollar denominated) dated as of June 6, 2001, among Fresenius Medical Care AG, FMC Trust Finance S.à.r.l. Luxembourg-III, State Street Bank and Trust Company, as Trustee, and the Subsidiary Guarantors named therein (Incorporated by reference to Exhibit No. 4.46 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.22 Senior Subordinated Indenture (Euro denominated) dated as of June 15, 2001, among Fresenius Medical Care AG, FMC Trust Finance S.à.r.l. Luxembourg-III, State Street Bank and Trust Company, as Trustee, and the Subsidiary Guarantors named therein (Incorporated by reference to Exhibit No. 4.47 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.23 Guarantee Agreement dated as of June 6, 2001 between Fresenius Medical Care AG and State Street Bank and Trust Company as Trustee, with respect to Fresenius Medical Care Capital Trust IV (Incorporated by reference to Exhibit No. 4.48 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.24 Guarantee Agreement dated as of June 15, 2001 between Fresenius Medical Care AG and State Street Bank and Trust Company as Trustee, with respect to Fresenius Medical Care Capital Trust V (Incorporated by reference to Exhibit No. 4.49 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.25 Agreement as to Expenses and Liabilities between Fresenius Medical Care AG and Fresenius Medical Care Capital Trust IV dated as of June 6, 2001 (Incorporated by reference to Exhibit No. 4.50 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.26 Agreement as to Expenses and Liabilities between Fresenius Medical Care AG and Fresenius Medical Care Capital Trust V dated as of June 15, 2001 (Incorporated by reference to Exhibit No. 4.51 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558).

2.27 Receivables Purchase Agreement dated August 28, 1997 between National Medical Care, Inc. and NMC Funding Corporation. (Incorporated by reference to Exhibit 10.3 to FMCH's Quarterly Report on Form 10-Q, for the three months ended September 30, 1997, filed November 4, 1997).

2.28 Amendment dated as of September 28, 1998 to the Receivables Purchase Agreement dated as of August 28, 1997, by and between NMC Funding Corporation, as Purchaser and National Medical Care, Inc., as Seller. (Incorporated by reference to Exhibit 10.1 to FMCH's Quarterly Report on Form 10-Q, for the three months ended September 30, 1998, filed November 12, 1998).

2.29 Third Amended and Restated Transfer and Administrative agreement dated as of October 23, 2003 among NMC Funding Corporation, National Medical Care, Inc., Paradigm Funding LLC, Asset One Securitization, LLC, Liberty Street Funding Corp., Giro Multifunding Corporation, and the Bank Investors listed therein, and WestLB AG, New York Branch, as administrative agent and agent (filed herewith).

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2.30 Amended and Restated Credit Agreement dated as of February 21, 2003 among Fresenius Medical Care AG and Fresenius Medical Care Holdings, Inc., as borrowers and guarantors, the direct and indirect subsidiaries of Fresenius Medical Care AG named therein as additional borrowers and guarantors, Bank of America N.A., as Administrative Agent, Credit Suisse First Boston, acting through its Cayman Islands Branch, and Dresdner Bank AG New York and Grand Cayman Branches, as Co-Documentation Agents, JPMorgan Chase Bank and The Bank of Nova Scotia, as Co-Syndication Agents and the Lenders party thereto (incorporated by reference to Exhibit No. 4.1 to the Form 10-Q of Fresenius Medical Care Holdings, Inc. for the three months ended March 31, 2003 filed May 15, 2003).⁽¹⁾

2.31 Amendment No. 1 dated as of August 22, 2003 to the Amended and Restated Credit Agreement dated as of February 21, 2003 among Fresenius Medical Care AG and Fresenius Medical Care Holdings, Inc., as borrowers and guarantors, the direct and indirect subsidiaries of Fresenius Medical Care AG named therein as additional borrowers and guarantors, Bank of America N.A., as Administrative Agent, Credit Suisse First Boston, acting through its Cayman Islands Branch, and Dresdner Bank AG New York and Grand Cayman Branches, as Co-Documentation Agents, JPMorgan Chase Bank and The Bank of Nova Scotia, as Co-Syndication Agents and the Lenders party thereto (incorporated by reference to Exhibit 4.2 to the Form 10-Q of Fresenius Medical Care Holdings, Inc. for the three month period ended September 30, 2003 filed November 13, 2003).⁽¹⁾

4.1 Agreement and Plan of Reorganization dated as of February 4, 1996 between W.R. Grace & Co. and Fresenius AG. (Incorporated by reference to Appendix A to the Joint Proxy Statement-Prospectus of Fresenius Medical Care AG, W.R. Grace & Co. and Fresenius USA, Inc., dated August 2, 1996).

4.2 Distribution Agreement by and among W.R. Grace & Co., W.R., Grace & Co. Conn. and Fresenius AG dated as of February 4, 1996. (Incorporated by reference to Appendix A to the Joint Proxy Statement-Prospectus of Fresenius Medical Care AG, W.R. Grace & Co. and Fresenius USA, Inc., dated August 2, 1996).

4.3 Contribution Agreement by and among Fresenius AG, Sterilpharma GmbH and W.R. Grace & Co. Conn. dated February 4, 1996. (Incorporated by reference to Appendix E to the Joint Proxy Statement-Prospectus of Fresenius Medical Care AG, W.R. Grace & Co. and Fresenius USA, Inc., dated August 2, 1996).

4.4 Post-Closing Covenants Agreement dated September 27, 1996 among Fresenius AG, W.R. Grace & Co., W.R. Grace & Co. Conn., and Fresenius Medical Care AG. (Incorporated by reference to Exhibit 10.11 to the Registrant's Registration Statement on Form F-1, filed on November 4, 1996).

4.5 Lease Agreement for Office Buildings dated September 30, 1996 by and between Fresenius AG and Fresenius Medical Care Deutschland GmbH. (Incorporated by reference to Exhibit 10.3 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.6 Lease Agreement for Manufacturing Facilities dated September 30, 1996 by and between Fresenius Immobilien-Verwaltungs-GmbH & Co. Objekt Schweinfurt KG and Fresenius Medical Care Deutschland GmbH. (Incorporated by reference to Exhibit 10.4 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.7 Lease Agreement for Manufacturing Facilities dated September 30, 1996 by and between Fresenius AG and Fresenius Medical Care Deutschland GmbH. (Incorporated by reference to Exhibit 10.5 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.8 Transition Services Agreement dated September 27, 1996 by and between Fresenius AG and Fresenius Medical Care. (Incorporated by reference to Exhibit 10.6 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.9 Forms of Supply Agreements between subsidiaries of Fresenius AG and subsidiaries of Fresenius Medical Care AG. (Incorporated by reference to Exhibit 10.7 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.10 Trademark License Agreement dated September 27, 1996 by and between Fresenius AG and Fresenius Medical Care AG. (Incorporated by reference to Exhibit 10.8 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

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4.11 Technology License Agreement (Biofine) dated September 27, 1996 by and between Fresenius AG and Fresenius Medical Care AG. (Incorporated by reference to Exhibit 10.9 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.12 Cross-License Agreement dated September 27, 1996 by and between Fresenius AG and Fresenius Medical Care AG. (Incorporated by reference to Exhibit 10.10 to the Registrant's Registration Statement on Form F-1, Registration No. 333-05922, filed November 16, 1996).

4.13 Lease Agreement for Office Buildings dated September 30, 1996 by and between Fresenius AG and Fresenius Medical Care Deutschland GmbH (Daimler Str.). (Incorporated by reference to Exhibit 2.8 to the Registrant's Annual Report on Form 20-F for the year ended December 31, 1996, filed April 7, 1997).

4.14 Fresenius Medical Care AG 1996 Stock Incentive Plan, (incorporated by reference to the Registrant's Registration Statement on Form S-8, dated October 1, 1996).

4.15 Fresenius Medical Care AG Rollover Stock Option Plan (Incorporated by reference to the Registrant's Registration Statement on Form S-8, dated September 30, 1996).

4.16 Fresenius Medical Care AG 1998 Stock Incentive Plan adopted effective as of April 6, 1998. (Incorporated by reference to Exhibit 4.8 to the Registrant's Report on Form 6-K for the three months ended March 31, 1998, filed May 14, 1998).

4.17 Fresenius Medical Care AG Stock Option Plan of June 10, 1998 (for non-North American employees). (Incorporated by reference to Exhibit 1.2 to the Registrant's Annual Report on Form 20-F, for the year ended December 31, 1998, filed March 24, 1999).

4.18 Fresenius Medical Care Aktiengesellschaft 2001 International Stock Incentive Plan (Incorporated by reference to Exhibit No. 10.17 to the Registration Statement on Form F-4 of Fresenius Medical Care AG et al filed August 2, 2001, Registration No. 333-66558

4.19 Product Purchase Agreement, effective January 1, 2004 between Amgen, Inc. and National Medical Care, Inc. (filed herewith).⁽¹⁾

4.20 Corporate Integrity Agreement dated January 18, 2000 between FMCH and Office of the Inspector General of the Department of Health and Human Services. (Incorporated by reference to Exhibit 10.1 to FMCH's Current Report on Form 8-K dated January 21, 2000).

4.21 Settlement Agreement dated as of February 6, 2003 by and among Fresenius Medical Care AG, Fresenius Medical Care Holdings, National Medical Care, Inc., the Official Committee of Asbestos Personal Injury Claimants, and the Official Committee of Asbestos Property Damage Claimants of W.R.Grace & Co. (incorporated by reference to Exhibit No. 10.18 on Form 10-K of Fresenius Medical Care Holdings, Inc. for the year ended December 31, 2002 filed March 17, 2002).

8.1 List of Significant Subsidiaries. Our significant subsidiaries are identified in Item 4.C. Information on the Company Organizational Structure.

11.1 Code of Business Conduct for Fresenius Medical Care AG, last revised in December, 2003 (filed herewith).

12.1 Certification of Chief Executive Officer Pursuant to Section 302 of the Sarbanes-Oxley Act of 2002 (filed herewith).

12.2 Certification of Chief Financial Officer Pursuant to Section 302 of the Sarbanes-Oxley Act of 2002 (filed herewith).

13.1 Certification of Chief Executive Officer and Chief Financial Officer Pursuant to 18 U.S.C. Section 1350, as Adopted Pursuant to Section 906 of the Sarbanes-Oxley Act of 2002 (filed herewith). (This Exhibit is furnished herewith, but not deemed filed for purposes of Section 18 of the Securities Exchange Act of 1934, as amended, or otherwise subject to liability under that section. Such certification will not be deemed to

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be incorporated by reference into any filing under the Securities Act or the Exchange Act, except to the extent that we explicitly incorporate it by reference.)

14.1 Consent of KPMG Deutsche Treuhand-Gesellschaft Aktiengesellschaft Wirtschaftsprüfungsgesellschaft (filed herewith).

14.2 Pages 114-116 from the final prospectus of Fresenius Medical Care AG dated July 20, 2000, consisting of the information under the heading DESCRIPTION OF THE POOLING AGREEMENTS (incorporated by reference to Exhibit No. 10.2 on the registrants Form 20-F for the year ended December 31, 2002 filed March 18, 2003).

(1) Confidential treatment has been requested as to certain portions of this document in accordance with the applicable rules of the Securities and Exchange Commission.

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SIGNATURES

The Registrant hereby certifies that it meets all of the requirements for filing on Form 20-F and that it has duly caused and authorized the undersigned to sign this annual report on its behalf.

DATE: March 2, 2004

FRESENIUS MEDICAL CARE
AKTIENGESELLSCHAFT

By: /s/ DR. BEN LIPPS

Name: Dr. Ben Lipps
Title: Chairman of the Management Board

By: /s/ LAWRENCE ROSEN

Name: Lawrence Rosen
Title: Chief Financial Officer

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INDEPENDENT AUDITORS REPORT

To the Shareholders

Fresenius Medical Care Aktiengesellschaft

Hof an der Saale, Germany:

We have audited the accompanying consolidated balance sheets of Fresenius Medical Care Aktiengesellschaft and subsidiaries (the Company) as of December 31, 2003 and 2002 and the related consolidated statements of operations, cash flows and shareholders' equity for each of the years in the three-year period ended December 31, 2003. In connection with our audits of the consolidated financial statements, we have also audited the financial statement schedule as listed in the accompanying index. These consolidated financial statements and the financial statement schedule are the responsibility of the Company's management. Our responsibility is to express an opinion on these consolidated financial statements and the financial statement schedule based on our audits.

We conducted our audits in accordance with auditing standards generally accepted in the United States of America. Those standards require that we plan and perform the audit to obtain reasonable assurance about whether the consolidated financial statements are free of material misstatement. An audit includes examining, on a test basis, evidence supporting the amounts and disclosures in the financial statements. An audit also includes assessing the accounting principles used and significant estimates made by management, as well as evaluating the overall financial statement presentation. We believe that our audits provide a reasonable basis for our opinion.

In our opinion, the consolidated financial statements referred to above present fairly, in all material respects, the financial position of the Company as of December 31, 2003 and 2002, and the results of its operations and its cash flows for each of the years in the three-year period ended December 31, 2003, in conformity with accounting principles generally accepted in the United States of America. Also in our opinion, the related financial statement schedule, when considered in relation to the consolidated financial statements taken as a whole, presents fairly, in all material respects, the information set forth therein.

As discussed in Note 1 to the consolidated financial statement, effective January 1, 2002, the Company adopted Statement of Financial Accounting Standards No. 142, *Goodwill and Other Intangible Assets*.

Frankfurt am Main, Germany

February 10, 2004

/s/ KPMG

Deutsche Treuhand-Gesellschaft
Aktiengesellschaft
Wirtschaftsprüfungsgesellschaft

Table of Contents**Fresenius Medical Care AG****Consolidated Statements of Operations**

For the years ended December 31, 2003, 2002 and 2001
(in thousands, except share data)

	2003	2002	2001
Net revenue:			
Dialysis Care	\$ 3,978,344	\$ 3,708,903	\$ 3,557,234
Dialysis Products	1,549,165	1,375,194	1,302,084
	<u>5,527,509</u>	<u>5,084,097</u>	<u>4,859,318</u>
Costs of revenue:			
Dialysis Care	2,871,592	2,713,341	2,521,075
Dialysis Products	827,014	714,736	699,123
	<u>3,698,606</u>	<u>3,428,077</u>	<u>3,220,198</u>
Gross profit	1,828,903	1,656,020	1,639,120
Operating expenses:			
Selling, general and administrative	1,021,781	913,620	966,044
Research and development	49,687	47,433	35,700
Special charge for legal matters			258,159
	<u>1,071,468</u>	<u>961,053</u>	<u>1,260,003</u>
Operating income	757,435	694,967	379,217
Other (income) expense:			
Interest income	(19,089)	(18,053)	(14,305)
Interest expense	230,848	244,570	237,234
	<u>211,759</u>	<u>262,623</u>	<u>252,929</u>
Income before income taxes and minority interest	545,676	468,450	156,288
Income tax expense	212,714	175,074	91,202
Minority interest	1,782	3,586	1,732
	<u>214,500</u>	<u>178,650</u>	<u>104,134</u>
Net income	<u>\$ 331,180</u>	<u>\$ 289,790</u>	<u>\$ 63,354</u>
Basic income per Ordinary share	<u>\$ 3.42</u>	<u>\$ 3.00</u>	<u>\$ 0.65</u>
Fully diluted income per Ordinary share	<u>\$ 3.42</u>	<u>\$ 3.00</u>	<u>\$ 0.64</u>
Basic income per Preference share	<u>\$ 3.49</u>	<u>\$ 3.06</u>	<u>\$ 0.70</u>
Fully diluted income per Preference share	<u>\$ 3.49</u>	<u>\$ 3.06</u>	<u>\$ 0.69</u>

See accompanying notes to consolidated financial statements.

Table of Contents**Fresenius Medical Care AG****Consolidated Balance Sheets**

At December 31, 2003 and 2002
(in thousands, except share data)

	<u>2003</u>	<u>2002</u>
Assets		
Current assets:		
Cash and cash equivalents	\$ 48,427	\$ 64,793
Trade accounts receivable, less allowance for doubtful accounts of \$166,385 in 2003 and \$159,763 in 2002	1,229,503	914,302
Accounts receivable from related parties	50,456	41,332
Inventories	444,738	372,222
Prepaid expenses and other current assets	253,365	239,172
Deferred taxes	179,639	189,879
	<u>2,206,128</u>	<u>1,821,700</u>
Total current assets	2,206,128	1,821,700
Property, plant and equipment, net	1,089,146	917,868
Intangible assets	582,103	550,321
Goodwill	3,288,348	3,192,651
Deferred taxes	35,541	35,741
Other assets	302,054	261,668
	<u>7,503,320</u>	<u>6,779,949</u>
Total assets	\$ 7,503,320	\$ 6,779,949
Liabilities and shareholders' equity		
Current liabilities:		
Accounts payable	\$ 177,824	\$ 185,949
Accounts payable to related parties	128,703	98,992
Accrued expenses and other current liabilities	553,830	469,228
Accrual for special charge for legal matters	138,154	191,130
Short-term borrowings	89,417	124,964
Short-term borrowings from related parties	30,000	6,000
Current portion of long-term debt and capital lease obligations	90,365	22,394
Income tax payable	178,111	178,690
Deferred taxes	26,077	18,027
	<u>1,412,481</u>	<u>1,295,374</u>
Total current liabilities	1,412,481	1,295,374
Long-term debt and capital lease obligations, less current portion	1,111,624	1,089,210
Other liabilities	128,615	154,859
Pension liabilities	100,052	96,152
Deferred taxes	250,446	169,372
Company-obligated mandatorily redeemable preferred securities of subsidiary Fresenius Medical Care Capital Trusts holding solely		
Company-guaranteed debentures of subsidiaries	1,242,317	1,145,281
Minority interest	14,105	22,522
	<u>4,259,640</u>	<u>3,972,770</u>
Total liabilities	4,259,640	3,972,770
Shareholders' equity:		
Preference shares, no par, 2.56 nominal value, 53,597,700 shares authorized, 26,213,979 issued and outstanding	69,616	69,540
Ordinary shares, no par, 2.56 nominal value, 70,000,000 shares authorized, issued and outstanding	229,494	229,494
Additional paid-in capital	2,741,362	2,736,913

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Retained earnings	378,014	154,595
Accumulated other comprehensive loss	(174,806)	(383,363)
	<u> </u>	<u> </u>
Total shareholders' equity	3,243,680	2,807,179
	<u> </u>	<u> </u>
Total liabilities and shareholders' equity	\$7,503,320	\$6,779,949
	<u> </u>	<u> </u>

See accompanying notes to consolidated financial statements.

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Table of Contents**Fresenius Medical Care AG****Consolidated Statements of Cash Flows**

For the years ended December 31, 2003, 2002 and 2001
(in thousands)

	2003	2002	2001
Operating Activities:			
Net income	\$ 331,180	\$ 289,790	\$ 63,354
Adjustments to reconcile net income to cash and cash equivalents provided by (used in) operating activities:			
Depreciation and amortization	216,377	210,555	323,503
Loss on early redemption of trust preferred securities, net of tax		11,777	
Change in deferred taxes, net	91,312	58,449	(46,401)
(Gain) loss on sale of fixed assets	(50)	690	1,010
Compensation expense related to stock options	1,456	1,126	1,153
Cash inflow from Hedging	131,654	24,542	
Changes in assets and liabilities, net of amounts from businesses acquired:			
Trade accounts receivable, net	53,563	(13,124)	(117,093)
Inventories	(22,993)	(6,519)	(30,201)
Prepaid expenses, other current and non-current assets	60,155	17,670	(28,462)
Accounts receivable from/payable to related parties	7,199	3,228	8,854
Accounts payable, accrued expenses and other current and non-current liabilities	(92,316)	(42,518)	183,992
Income tax payable	(23,518)	(5,748)	64,539
Net cash provided by operating activities	<u>754,019</u>	<u>549,918</u>	<u>424,248</u>
Investing Activities:			
Purchases of property, plant and equipment	(291,260)	(239,160)	(275,225)
Proceeds from sale of property, plant and equipment	14,826	37,783	24,195
Acquisitions and investments, net of cash acquired	(92,190)	(79,835)	(216,711)
Net cash used in investing activities	<u>(368,624)</u>	<u>(281,212)</u>	<u>(467,741)</u>
Financing Activities:			
Proceeds from short-term borrowings	102,678	88,639	117,896
Repayments of short-term borrowings	(153,911)	(68,255)	(140,420)
Proceeds from short-term borrowings from related parties	94,787	49,120	20,588
Repayments of short-term borrowings from related parties	(70,787)	(58,125)	(223,566)
Proceeds from long-term debt	982,825	417,098	465,906
Principal payments of long-term debt and capital lease obligations	(968,888)	(246,566)	(517,877)
Payments on obligation related to 1999 Settlement			(85,920)
Proceeds from issuance of trust preferred securities			470,598
Redemption of trust preferred securities		(376,200)	
(Decrease) increase of accounts receivable securitization program	(287,251)	3,249	(3,464)
Proceeds from exercise of stock options	1,600	550	6,391
Dividends paid	(107,761)	(76,743)	(65,782)
Redemption of Series D Preferred Stock of subsidiary	(8,906)		
Change in other minority interest	(266)	2,095	(853)
Net cash (used in) provided by financing activities	<u>(415,880)</u>	<u>(265,138)</u>	<u>43,497</u>

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Effect of exchange rate changes on cash and cash equivalents	14,119	(347)	(3,009)
Cash and Cash Equivalents:			
Net (decrease) increase in cash and cash equivalents	(16,366)	3,221	(3,005)
Cash and cash equivalents at beginning of period	64,793	61,572	64,577
Cash and cash equivalents at end of period	\$ 48,427	\$ 64,793	\$ 61,572

See accompanying notes to consolidated financial statements.

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Table of Contents**Fresenius Medical Care AG****Consolidated Statements of Shareholders' Equity**

For the years ended December 31, 2003, 2002 and 2001
(in thousands, except share data)

	Preference Shares		Ordinary Shares		Accumulated other comprehensive loss					Total
	Number of shares	No par value	Number of shares	No par value	Additional paid in capital	Retained earnings (deficit)	Foreign currency translation	Cash Flow Hedges	Minimum Pension Liability	
Balance at December 31, 2000	23,765,093	\$ 63,644	70,000,000	\$ 229,494	\$ 2,634,606	\$ (56,024)	\$ (192,970)	\$	\$	\$ 2,678,750
Issuance of Preference shares	2,250,000	5,498			93,485					98,983
Proceeds from exercise of options	161,415	371			6,020					6,391
Compensation expense related to stock options					1,153					1,153
Dividends paid						(65,782)				(65,782)
Comprehensive loss										
Net income						63,354				63,354
Other comprehensive loss related to cash flow hedges								(50,683)		(50,683)
Foreign currency translation adjustment							(115,422)			(115,422)
Comprehensive loss										(102,751)
Balance at December 31, 2001	26,176,508	\$ 69,512	70,000,000	\$ 229,494	\$ 2,735,265	\$ (58,452)	\$ (308,392)	\$ (50,683)	\$	\$ 2,616,744
Proceeds from exercise of options	12,067	28			522					550
Compensation expense related to stock options					1,126					1,126
Dividends paid						(76,743)				(76,743)
Comprehensive income										
Net income						289,790				289,790
Other comprehensive income related to:										
Cash flow hedges								33,501		33,501
Foreign currency translation adjustment							(38,432)			(38,432)
Minimum pension liability									(19,357)	(19,357)
Comprehensive income										265,502
Balance at December 31, 2002	26,188,575	\$ 69,540	70,000,000	\$ 229,494	\$ 2,736,913	\$ 154,595	\$ (346,824)	\$ (17,182)	\$ (19,357)	\$ 2,807,179
	25,404	76			1,524					1,600

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Proceeds from exercise of options										
Compensation expense related to stock options				1,456						1,456
Dividends paid					(107,761)					(107,761)
Transaction under common control with Fresenius AG				1,469						1,469
Comprehensive income										
Net income					331,180					331,180
Other comprehensive income related to:										
Cash flow hedges								22,029		22,029
Foreign currency translation adjustment						200,578				200,578
Minimum Pension Liability									(14,050)	(14,050)
Comprehensive income										539,737
Balance at December 31, 2003	26,213,979	\$ 69,616	70,000,000	\$ 229,494	\$ 2,741,362	\$ 378,014	\$ (146,246)	\$ 4,847	\$ (33,407)	\$ 3,243,680

See accompanying notes to consolidated financial statements.

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Fresenius Medical Care AG

Notes to Consolidated Financial Statements

(in thousands, except share data)

1. The Company and Summary of Significant Accounting Policies

Fresenius Medical Care AG and subsidiaries (FMS or the Company), is an integrated provider of kidney dialysis products and dialysis care. FMS was created by conversion of Sterilpharma GmbH, a limited liability company incorporated in 1975, into a stock corporation (Aktiengesellschaft). The resolutions for this conversion were adopted by a shareholder meeting on April 17, 1996. On September 30, 1996, FMS initiated a series of transactions to consummate an Agreement and Plan of Reorganization entered into on February 4, 1996 by Fresenius AG and W.R. Grace & Co. (W.R. Grace). Pursuant to that Agreement, Fresenius AG contributed Fresenius Worldwide Dialysis or FWD, its global dialysis business, including its controlling interest in Fresenius USA, Inc. (FUSA), in exchange for FMS Ordinary shares. Thereafter, FMS, in exchange for Ordinary shares, acquired: (i) all of the outstanding Common stock of W.R. Grace, whose sole business at the time of the transaction consisted of National Medical Care, Inc. (NMC), its global dialysis business; and (ii) the publicly-held minority interest of FUSA.

Basis of Presentation

The accompanying consolidated financial statements have been prepared in accordance with accounting principles generally accepted in the United States of America (U.S. GAAP).

Summary of Significant Accounting Policies

a) Principles of Consolidation

The consolidated financial statements include all material companies in which the Company has legal or effective control. The equity method of accounting is used for investments in associated companies (20% to 50% owned). All significant intercompany transactions and balances have been eliminated.

b) Classifications

Certain items in prior years consolidated financial statements may have been reclassified to conform with the current year s presentation. Net operating results have not been affected by the reclassifications.

c) Cash and Cash Equivalents

Cash and cash equivalents represent cash and certificates of deposit with original maturity dates of three months or less at origination.

d) Allowance for Doubtful Accounts

Estimates for the allowances for accounts receivable from the dialysis service business are mainly based on past collection history. Specifically, the allowances for the North American services division are based on an analysis of collection experience, recognizing the differences between payors and aging of accounts receivable. From time to time, accounts receivable are reviewed for changes from the historic collection experience to ensure the appropriateness of the allowances. The allowances in the international segment and the products business are based on estimates and consider various factors, including aging, creditor and past collection history.

e) Inventories

Inventories are stated at the lower of cost (determined by using the average or first-in, first-out method) or market value.

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f) Property, Plant and Equipment

Property, plant, and equipment are stated at cost less accumulated depreciation. Significant improvements are capitalized; repairs and maintenance costs that do not extend the useful lives of the assets are charged to expense as incurred. Property and equipment under capital leases are stated at the present value of future minimum lease payments at the inception of the lease, less accumulated depreciation. Depreciation on property, plant and equipment is calculated using the straight-line method over the estimated useful lives of the assets ranging from 5 to 50 years for buildings and improvements with a weighted average life of 12 years and 3 to 15 years for machinery and equipment with a weighted average life of 8 years. Equipment held under capital leases and leasehold improvements are amortized using the straight-line method over the shorter of the lease term or the estimated useful life of the asset. The Company capitalizes interest on borrowed funds during construction periods. Interest capitalized during 2003, 2002, and 2001 was \$920, \$3,248 and \$3,532, respectively.

g) Goodwill and Intangible Assets

In accordance with SFAS 141, *Business Combinations*, the Company applies the purchase method for all business combinations. Intangible assets acquired in a purchase method business combination are recognized and reported apart from goodwill, pursuant to the criteria specified by SFAS No. 141.

The Company adopted SFAS No. 142, *Goodwill and Other Intangible Assets*. Pursuant to SFAS No. 142, intangible assets with finite useful lives are amortized over their respective estimated useful lives to their estimated residual values, and reviewed for impairment in accordance with SFAS No. 144, *Accounting for the Impairment or Disposal of Long-Lived Assets* (see m) Impairment).

As of January 1, 2002, in accordance with SFAS No. 142, goodwill and identifiable intangibles with indefinite lives are no longer amortized, but tested annually for impairment. The Company identified trade names and management contracts as intangible assets with indefinite useful lives.

To evaluate the recoverability of goodwill, the Company identified its reporting units and determined the carrying value of each reporting unit by assigning the assets and liabilities, including the existing goodwill and intangible assets, to those reporting units. At least once a year the Company compares the fair value of each reporting unit to the reporting unit's carrying amount. Fair value is determined using a discounted cash flow approach. In the case that the fair value of the reporting unit is less than its book value, a second step is performed which compares the fair value of the reporting unit's goodwill to the carrying value of its goodwill. If the fair value of the goodwill is less than the book value, the difference is recorded as an impairment.

To evaluate the recoverability of intangible assets with indefinite useful lives, the Company compares the fair values of intangible assets with their carrying value. An intangible asset's fair value is determined using a discounted cash flow approach and other appropriate methods.

h) Derivative Financial Instruments

The Company adopted SFAS No. 133, *Accounting for Derivative Instruments and Hedging Activities* as amended by SFAS No. 138 and SFAS 149. The Company utilizes derivative financial instruments including forward currency contracts and interest rate swaps. SFAS No. 133 requires all derivatives to be recognized as assets or liabilities at fair value.

Changes in the fair value of foreign currency forward contracts designated and qualifying as effective cash flow hedges of forecasted transactions are reported in accumulated other comprehensive income. These amounts are subsequently reclassified into earnings as a component of the forecasted transaction in the same period as the forecasted transaction affects earnings.

Changes in the fair value of interest rate swaps that are designated as cash flow hedges and effectively convert variable interest payments into fixed interest payments are deferred in accumulated other comprehensive income. The interest rate agreements are accounted for on an accrual basis, i.e. the interest payable and the interest rate receivable under the terms of the swaps are accrued and recorded as an adjustment to the interest or related expense of the designated liability or obligation.

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Amounts due from and payable to the counterparties of interest rate swaps are recorded on an accrual basis at each reporting date at amounts computed by reference to the respective interest rate swap contract. Realized gains and losses that occur from the early termination or expiration of contracts are deferred and recorded in income over the remaining period of the original swap agreement if the corresponding debt is still outstanding. Gains and losses arising from interest differential on contracts that hedge specific borrowings are recorded as a component of interest expense over the life of the contract. In the event the hedged asset or liability is terminated, sold, or otherwise disposed of, the gain or loss on the interest rate swap would be matched with the offsetting gain or loss of the related item (see Note 21).

i) Foreign Currency Translation

For purposes of these consolidated financial statements, the U.S. dollar is the reporting currency. The Company follows the provisions of SFAS No. 52, *Foreign Currency Translation*. Substantially all assets and liabilities of the parent company and all non-U.S. subsidiaries are translated at year-end exchange rates, while revenues and expenses are translated at exchange rates prevailing during the year. Adjustments for foreign currency translation fluctuations are excluded from net earnings and are reported in accumulated other comprehensive income. In addition, the translation adjustments of certain intercompany borrowings, which are considered foreign equity investments, are reported in accumulated other comprehensive income.

j) Revenue Recognition Policy

Dialysis care revenues are recognized on the date services and related products are provided and the payor is obligated to pay at amounts estimated to be received under reimbursement arrangements with third party payors. Medicare and Medicaid in North America and programs involving other government payors in the international segment are billed at pre-determined rates per treatment that are established by statute or regulation. Most non-governmental payors, including contracted managed care payors, are billed at our standard rates for services net of contractual allowances to reflect the estimated amounts to be received under reimbursement arrangements with these payors.

Dialysis product revenues are recognized when title to the product passes to the customers either at the time of shipment, upon receipt by the customer or upon any other terms that clearly define passage of title. As product returns are not typical, no return allowances are established. In the event a return is required, the appropriate reductions to sales, accounts receivables and cost of sales are made.

A minor portion of International product revenue are generated from arrangements which give the customer, typically a health care provider, the right to use dialysis machines. In the same contract the customer agrees to purchase the related treatment disposables at a price marked up from the standard price list. FMS does not recognize revenue for the delivery of the dialysis machine but recognizes revenue, including the mark-up on the sale of disposables.

k) Research and Development expenses

Research and development expenses are expensed as incurred.

l) Income Taxes

In accordance with SFAS No. 109, *Accounting for Income Taxes*, deferred tax assets and liabilities are recognized for the future consequences attributable to differences between the financial statement carrying amounts of existing assets and liabilities and their respective tax basis. Deferred tax assets and liabilities are measured using enacted tax rates expected to apply to taxable income in the years in which those temporary differences are expected to be recovered or settled. A valuation allowance is recorded to reduce the carrying amount of the deferred tax assets unless it is more likely than not that such assets will be realized. (see Note 18)

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m) Impairment

The Company calculates extraordinary amortization in accordance with SFAS No. 144, *Accounting for the Impairment or Disposal of Long-Lived Assets*. The Company reviews the carrying value of its long-lived assets or asset groups with definite useful lives to be held and used for impairment whenever events or changes in circumstances indicate that the carrying value of these assets may not be recoverable. Recoverability of these assets is measured by a comparison of the carrying value of an asset to the future net cash flow directly associated with the asset. If assets are considered to be impaired, the impairment recognized is the amount by which the carrying value exceeds the fair value of the asset. The Company uses various valuation factors, including market prices and present value techniques to assess fair value.

In accordance with SFAS No. 144, long-lived assets to be disposed of by sale are reported at the lower of carrying value or fair value less cost to sell and depreciation is ceased. Long-lived assets to be disposed of other than by sale are considered to be held and used until disposal.

n) Debt Issuance Costs

Costs related to the issuance of debt are amortized over the term of the related obligation.

o) Self-Insurance Programs

The Company's largest subsidiary is partially self-insured for professional, product and general liability, auto and worker's compensation claims under which the company assumes responsibility for incurred claims up to predetermined amounts above which third party insurance applies. Reported balances for the year include estimates of the anticipated expense for claims incurred (both reported and incurred but not reported) based on historical experience and existing claim activity. This experience includes both the rate of claims incidence (number) and claim severity (cost) and is combined with individual claim expectations to estimate the reported amounts.

p) Use of Estimates

The preparation of consolidated financial statements in conformity with U.S. GAAP requires management to make estimates and assumptions that affect the reported amounts of assets and liabilities and disclosure of contingent assets and liabilities at the date of the consolidated financial statements and the reported amounts of revenues and expenses during the reporting period. Actual results could differ from those estimates.

q) Concentration of Credit Risk

The Company is engaged in the manufacture and sale of products for all forms of kidney dialysis, principally to health care providers throughout the world, and in providing kidney dialysis treatment, clinical laboratory testing and other medical ancillary services. The Company performs ongoing evaluations of its customers' financial condition and, generally, requires no collateral.

Approximately 43%, 43% and 42% of the Company's worldwide revenues are paid by and subject to regulations under governmental health care programs, primarily Medicare and Medicaid, administered by the United States government in 2003, 2002, and 2001, respectively.

r) Earnings per Preference share and Ordinary share

Basic net income per Preference share and basic net income per Ordinary share for all years presented have been calculated using the two-class method required under U.S. GAAP based upon the weighted average number of Ordinary and Preference shares outstanding. Basic earnings per share are computed by dividing net income less preference amounts by the weighted average number of Ordinary shares and Preference shares outstanding during the year. Diluted earnings per share include the effect of all potentially dilutive Ordinary shares and Preference shares that would have been outstanding during the year.

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The awards granted under the Company's stock incentive plans (see Note 17), are potentially dilutive equity instruments.

s) Stock Option Plans

The Company accounts for its stock option plans using the intrinsic value method in accordance with the provisions of Accounting Principles Board (APB) Opinion No. 25, *Accounting for Stock Issued to Employees*, and related interpretations. As such, compensation expense is recorded only if the current market price of the underlying stock exceeds the exercise price on the measurement date. For stock incentive plans which are performance based, the Company recognizes compensation expense over the vesting periods, based on the then current market values of the underlying stock.

The following table illustrates the effect on net income and earnings per share if the Company had applied the fair value recognition provisions of SFAS No. 123, *Accounting for Stock-Based Compensation*, to stock based employee compensation.

	2003	2002	2001
Net income:			
As reported	\$ 331,180	\$ 289,790	\$ 63,354
Add: Stock-based employee compensation expense included in reported net income, net of related tax effects	1,456	1,126	1,153
Deduct: Total stock-based employee compensation expense determined under fair value method for all awards, net of related tax effects	(9,583)	(11,951)	(13,223)
Pro forma	\$ 323,053	\$ 278,965	\$ 51,284
Basic net income per:			
Ordinary share			
As reported	\$ 3.42	\$ 3.00	\$ 0.65
Pro forma	\$ 3.34	\$ 2.88	\$ 0.52
Preference share			
As reported	\$ 3.49	\$ 3.06	\$ 0.70
Pro forma	\$ 3.41	\$ 2.94	\$ 0.57
Fully diluted net income per:			
Ordinary share			
As reported	\$ 3.42	\$ 3.00	\$ 0.64
Pro forma	\$ 3.34	\$ 2.88	\$ 0.51
Preference share			
As reported	\$ 3.49	\$ 3.06	\$ 0.69
Pro forma	\$ 3.41	\$ 2.94	\$ 0.56

t) Recent Pronouncements and Accounting Changes

In August 2001, the FASB issued SFAS No. 143, *Accounting for Asset Retirement Obligations*. SFAS No. 143 requires that the fair value of a liability for an asset retirement obligation be recognized in the period in which it is incurred if a reasonable estimate of fair value can be made. The associated asset retirement costs are capitalized as part of the carrying amount of the long-lived asset. It applies to legal obligations associated with the retirement of long-lived assets that result from the acquisition, construction, development and/or normal operation of a long-lived asset. The Company adopted SFAS No. 143 as of January 1, 2003. The adoption of SFAS No. 143 did not have a material impact on the Company's financial statements.

In April 2002, the FASB issued SFAS No. 145, *Rescission of FASB Statements No. 4, 44, and 64, Amendment of FASB Statement No. 13, and Technical Corrections*. SFAS No. 145 rescinds SFAS No. 4, SFAS No. 64 related to classifications of gains and losses on debt extinguishments such that most debt extinguishment gains and losses will no longer be classified as extraordinary. SFAS No. 145 also amends SFAS No. 13, with

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respect to certain sale-leaseback transactions. The Company adopted SFAS No. 145 in regard to SFAS No. 4 on January 1, 2003. In the first quarter of 2002, the Company recorded an extraordinary loss of approximately \$11,800, net of taxes of approximately \$7,700, as a result of the early redemption of debt (see Note 13). This loss is no longer presented as an extraordinary loss upon the adoption of SFAS No. 145. The Company adopted the other provisions of SFAS No. 145 effective April 1, 2002.

In June 2002, the FASB issued SFAS No. 146, *Accounting for Costs Associated with Exit or Disposal Activities*. The standard requires companies to recognize costs associated with exit or disposal activities when liabilities are incurred. SFAS No. 146 replaces EITF Issue No. 94-3, *Liability Recognition for Certain Employee Termination Benefits and Other Costs to Exit an Activity (Including Certain Costs Incurred in a Restructuring)*. This statement is applied prospectively to exit or disposal activities initiated after December 31, 2002.

In November 2002, the Financial Accounting Standards Board issued FASB Interpretation No. 45 (FIN 45), *Guarantor's Accounting and Disclosure Requirements for Guarantees of Indebtedness of Others*. FIN 45 also requires the guarantor to recognize a liability for the non-contingent component of the guarantee, that is, the obligation to stand ready to perform in the event that special triggering events or conditions occur. The initial recognition and measurement provisions are applicable prospectively to guarantees issued or modified after December 31, 2002. FIN 45 also clarifies and expands the disclosure requirements related to guarantees, including product warranties. FIN 45 does not materially impact the Company's financial statements.

On April 3, 2003, the Financial Accounting Standards Board issued SFAS No. 149 *Amendment of Statement 133 on Derivative Instruments and Hedging Activities*. This Statement amends and clarifies financial accounting and reporting for derivative instruments, including certain derivative instruments embedded in other contracts (collectively referred to as derivatives) and for hedging activities under SFAS No. 133, *Accounting for Derivative Instruments and Hedging Activities*. This Statement is effective for contracts entered into or modified after June 30, 2003. This adoption did not have any impact on the Company's financial statements.

In May 2003, the Financial Accounting Standards Board issued SFAS No. 150 *Accounting for certain Financial Instruments with Characteristics of both Liabilities and Equity*. This Statement requires an issuer to classify certain financial instruments with the characteristics of both liabilities and equity as a liability (or asset in some circumstances) instead of equity. This Statement is effective for financial instruments entered into or modified after May 31, 2003 and otherwise effective at the beginning of the first interim period beginning after June 15, 2003. This adoption did not have any impact on the Company's financial statements.

In December 2003 the Financial Accounting Standards Board issued SFAS No. 132 (revised 2003) *Employers Disclosures about Pensions and Other Postretirement Benefits – an amendment of FASB Statements No. 87, 88 and 106*. This statement extends the publishing rules for pension liabilities according to SFAS No. 132. The accounting and valuation principles remain unchanged.

In December 2003, the Financial Accounting Standards Board issued FASB Interpretation No. 46R *Consolidation of Variable Interest Rate Entities (revised)* (FIN 46R) which addresses how a business enterprise should evaluate whether it has a controlling financial interest in an entity through means other than voting rights and accordingly should consolidate the entity. FIN 46R replaced FASB Interpretation No. 46 *Consolidation of Variable Interest Rate Entities* which was issued in January 2003. The Company is required to apply FIN 46R for special purpose entities as of December 31, 2003 and for all other Variable Interest Entities (VIEs) as of March 31, 2004. The Company is not involved with any special purpose entity which required initial consolidation as of December 31, 2003 and will apply FIN 46R on March 31, 2004 for all VIEs. The Company is party to various arrangements with certain dialysis clinics to provide management services, financing and product supply. Some of these clinics are variable interest entities. Under FIN 46R these clinics will be consolidated if the Company is the primary beneficiary. The Company also participates in a joint venture which is engaged in the perfusion business. The arrangements with the joint venture partner are such that it qualifies as a variable interest entity and the Company is the primary beneficiary. These variable interest entities generate approximately \$153,000 in annual revenue. This includes approximately \$14,000 related to variable interest entities in which the Company is not the primary beneficiary. The Company has investments, other long term assets and receivables of approximately \$42,000 which represent the Company's maximum exposure to loss as a result of its involvement with the variable interest entities.

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2. Special Charge for 1999 Settlement

On January 18, 2000, Fresenius Medical Care Holdings, Inc. (FMCH), NMC and certain other affiliated companies executed definitive agreements with the United States Government to settle (i) matters concerning violations of federal laws then under investigation and (ii) National Medical Care, Inc. 's claims with respect to outstanding Medicare receivables for intradialytic parenteral nutrition therapy (collectively, the Settlement).

In 2001, FMCH made a final payment to the U.S. Government of \$85,900 pursuant to the Settlement. In addition, FMCH received a final payment of \$5,200 in the first quarter of 2001 from the U.S. Government, related to FMCH 's claims for outstanding Medicare receivables. A letter of credit, purchased to secure the settlement payment obligation, was closed out with the last payment.

3. Special Charge for Legal Matters

In the fourth quarter of 2001, the Company recorded a \$258,159 (\$177,159 after tax) special charge to address 1996 merger-related legal matters, estimated liabilities and legal expenses arising in connection with the W.R. Grace Chapter 11 Proceedings and the cost of resolving pending litigation and other disputes with certain commercial insurers.

The Company accrued \$172,034 principally representing a provision for income taxes payable for the years prior to the 1996 merger for which the Company has been indemnified by W.R. Grace, but may ultimately be obligated to pay as a result of W.R. Grace 's Chapter 11 Proceedings. In addition, that amount included the estimated costs of defending the Company in all litigation arising out of W.R. Grace 's Chapter 11 Proceedings. During the second quarter of 2003, the court supervising W.R. Grace 's Chapter 11 Proceedings approved the definitive settlement agreement entered into among the Company, the committees representing asbestos creditors and W.R. Grace.

The Company included \$55,489 in the special charge to provide for settlement obligations, legal expenses and the resolution of disputed accounts receivable relating to various insurance companies. In November of 2003, the Company settled without litigation all claims raised by the final group of insurance companies who had contacted the Company concerning allegations of inappropriate billing practices and misrepresentations. The cost of the settlement will be charged against previously established accruals (See Note 20).

The remaining amount of the special charge (\$30,636 pretax) was accrued mainly for (i) assets and receivables that are impaired in connection with other legal matters and (ii) anticipated expenses associated with the continued defense and resolution of the legal matters.

Based on these developments, the Company has reduced its estimate for the settlement and related costs of the W.R. Grace Chapter 11 Proceedings by \$39,000. This reduction of the provision for the W.R. Grace matter has been applied to the other components of the special charge (i.e. reserves for settlement obligations and disputed accounts receivable from commercial insurers and other merger-related legal matters described in this note).

At December 31, 2003, there is a remaining balance of \$138,154 for the accrual for the special charge for legal matters. The Company believes that these reserves are adequate for the settlement of all matters described above. During the year ended December 31, 2003, \$52,976 in charges were applied against the accrued special charge for legal matters.

4. Related Party Transactions

a) Service Agreements

The Company is party to service agreements with Fresenius AG, the majority shareholder, and certain affiliates of Fresenius AG to receive services, including, but not limited to: administrative services, management information services, employee benefit administration, legal and environmental consultation and administration, insurance, central purchasing, tax services and treasury services. For the years 2003, 2002 and 2001, amounts charged by Fresenius AG to FMS under the terms of the agreements are \$26,172, \$23,012 and \$19,117, respectively. FMS also provides certain services to Fresenius AG and certain affiliates of Fresenius AG, including

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research and development, plant administration, patent administration and warehousing. FMS charged \$11,669, \$10,142 and \$6,134 for services rendered to Fresenius AG in 2003, 2002 and 2001, respectively.

Under operating lease agreements entered into with Fresenius AG, FMS paid Fresenius AG \$13,307, \$10,401 and \$9,239 during 2003, 2002 and 2001, respectively. The majority of the leases expire in 2006 with options for renewal.

b) Financing Provided by Fresenius AG

At December 31, 2003, the Company had short-term loans outstanding of \$30,000, which bore interest at an average rate of 1.165%. At December 31, 2002, the Company had short-term loans outstanding of \$6,000 which bore an average interest rate of 2.22%. Interest expense on these borrowings was, \$59, \$359 and \$6,887 for the years 2003, 2002 and 2001, respectively.

c) Products

During the years ended December 31, 2003, 2002 and 2001, the Company recognized sales of \$27,306, \$25,986 and \$24,063, respectively, to Fresenius AG and affiliates. During 2003, 2002 and 2001, the Company made purchases from Fresenius AG and affiliates in the amount of \$27,228, \$23,703 and \$19,703, respectively.

d) Acquisitions

During the second quarter of 2003 the Company acquired Fresenius AG's adsorber business for a purchase price of \$23,735, net of cash acquired. The adsorber business manufactures products used in the field of therapeutic apheresis. These therapies are similar to kidney dialysis treatment in that they consist of extracorporeal blood treatments. The acquisition was accounted for as a transaction under common control.

e) Other

During 1999, the Company granted to a member of the management board a five-year unsecured loan of \$2,000 with interest at 6.0% per annum. This loan was repaid in 2003.

A member of the Company's supervisory board is a partner in a law firm which provided services to the Company. The Company paid the law firm approximately \$483, \$292, and \$368 in 2003, 2002 and 2001, respectively.

A member of the Company's supervisory board was the chairman of the management board of a bank that served as one of two joint global coordinators of a public offering of Preference shares conducted by the Company in 2000. An affiliate of the bank purchased Preference shares in a private offering in 2000 and, in 2001, affiliates of the bank served as co-lead manager, and as an initial purchaser of a global offering of trust preferred securities. The Company paid fees and commissions of \$6,808 in total to the coordinators of the offering. The bank is also a lender and one of the Managing Agents under both the Company's original senior credit agreement and the new senior credit agreement dated February 21, 2003 (see Note 11).

The Vice Chairman of the Company's supervisory board is a member of the supervisory board of Fresenius AG, the majority holder of FMS's Ordinary shares. In May of 2003, the Chief Financial Officer of the Company resigned to assume the position of Chairman of the management board and CEO of Fresenius AG.

5. Acquisitions and Investments

The Company acquired certain health care and distribution facilities and other investments for a total consideration of \$101,250, \$87,876 and \$461,079 in 2003, 2002 and 2001, respectively. All acquisitions have been accounted for as purchase transactions and, accordingly, are included in the results of operations from the dates of acquisition. The excess of the total acquisition costs over the fair value of the tangible net assets acquired was approximately \$83,000, \$82,000 and \$367,000 for 2003, 2002 and 2001, respectively.

During the year ended December 31, 2003, the Company acquired certain health care and distribution facilities, including the adsorber business of Fresenius AG for \$92,190 in cash and assumed debt of \$9,060.

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In 2002, the Company's acquisitions principally involved individual dialysis clinics providing dialysis therapy. The consideration consisted of cash of \$79,835 and assumed debt of \$8,041.

In January 2001, the Company acquired Everest Healthcare Services Corporation (Everest) for \$365,000. The Everest operations acquired consist of approximately 70 clinics facilities providing dialysis therapy to approximately 6,800 patients in the eastern and central United States. Approximately \$99,000 was funded by the issuance of 2.25 million Fresenius Medical Care AG Preference shares to the Everest shareholders. The remaining purchase price was paid with \$131,000 cash and assumption of \$135,000 of debt. In 2001, aggregate consideration for all acquisitions consisted of cash of \$216,711, assumed debt of \$144,889 and \$99,479 in Preference shares.

6. Sale of Accounts Receivable

FMCH has an asset securitization facility (the accounts receivable facility) whereby certain receivables are sold to NMC Funding Corporation (NMC Funding), a wholly-owned subsidiary. NMC Funding then sells and assigns undivided percentage ownership interests in the receivables to certain bank investors. NMC Funding surrenders control over the ownership interests in the accounts receivables as a result of this sale. The ownership interests are removed from the consolidated balance sheets in accordance with SFAS No. 140. The retained interest in the accounts receivable is reflected on the face of the balance sheet net of uncollectable accounts to approximate fair value. The Company has a servicing obligation to act as collection agent on behalf of the bank investors. The accounts receivable facility was amended on October 23, 2003, extending its maturity to October 22, 2004.

At December 31, 2003 and 2002, \$157,998 and \$445,249, respectively, had been received pursuant to such sales and are reflected as reductions to accounts receivable. NMC Funding pays interest to the bank investors, calculated based on the commercial paper rates for the particular tranches selected. The effective interest rate was approximately 1.31% during 2003. Under the terms of the agreement, new interests in accounts receivable are sold without recourse as collections reduce previously sold accounts receivable. The costs related to such sales are expensed as incurred and recorded as interest expense and related financing costs.

7. Inventories

As of December 31, 2003 and 2002, inventories consisted of the following:

	2003	2002
Raw materials and purchased components	\$ 86,653	\$ 79,760
Work in process	33,778	26,233
Finished goods	244,355	196,830
Health care supplies	79,952	69,399
Inventories	<u>\$444,738</u>	<u>\$372,222</u>

Under the terms of certain unconditional purchase agreements, the Company is obligated to purchase approximately \$220,204 of materials, of which \$94,000 is committed at December 31, 2003 for 2004. The terms of these agreements run 1 to 6 years. Inventories as of December 31, 2003 include \$30,894 of Erythropoietin (EPO) which is supplied by a single source supplier in the United States. Delays, stoppages, or interruptions in the supply of EPO could adversely affect the operating results of the Company. Revenues from EPO accounted for approximately 23% of total revenue in the North America segment for both 2003 and 2002.

8. Property, Plant and Equipment

As of December 31, 2003 and 2002, property, plant and equipment consisted of the following:

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	2003	2002
Land and improvements	\$ 28,109	\$ 23,075
Buildings and improvements	694,327	586,251
Machinery and equipment	1,191,708	948,781
Machinery, equipment and rental equipment under capitalized leases	54,101	32,221
Construction in progress	58,509	62,257
	<u>2,026,754</u>	<u>1,652,585</u>
Accumulated depreciation	(937,608)	(734,717)
	<u>2,026,754</u>	<u>1,652,585</u>
Property, plant and equipment, net	<u>\$ 1,089,146</u>	<u>\$ 917,868</u>

Depreciation expense for property, plant and equipment amounted to \$180,952, \$158,126 and \$147,945 for the years ended December 31, 2003, 2002 and 2001, respectively.

Included in property, plant and equipment as of December 31, 2003 and 2002 were \$98,243 and \$89,754, respectively, of peritoneal dialysis cyclor machines which the Company leases to customers with end-stage renal disease on a month-to-month basis and hemodialysis machines which the Company leases to physicians under operating leases. Accumulated depreciation related to machinery, equipment and rental equipment under capital leases was \$29,654 and \$16,222 at December 31, 2003 and 2002, respectively.

9. Intangible Assets and Goodwill

As of December 31, 2003 and 2002, intangible assets other than goodwill consisted of the following:

	December 31, 2003		December 31, 2002		Average Useful Life
	Gross Carrying Amount	Accumulated Amortization	Gross Carrying Amount	Accumulated Amortization	
Amortizable Intangible Assets					
Patient relationships	\$ 258,408	\$(208,890)	\$ 249,069	\$(191,571)	14
Patents	18,178	(15,056)	14,395	(12,317)	27
Distribution rights	23,920	(9,548)	10,226	(5,886)	12
Other	170,320	(86,318)	155,317	(72,217)	11
	<u>\$ 470,826</u>	<u>\$(319,812)</u>	<u>\$ 429,007</u>	<u>\$(281,991)</u>	
Non-amortizable Intangible Assets					
Tradenname	221,720		220,249		
Management contracts	209,369		183,056		
	<u>\$ 431,089</u>		<u>\$ 403,305</u>		

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The related amortization expenses (including amortization for goodwill, tradename and management contracts in 2001) are as follows:

Aggregate Amortization Expense

For the year ended December 31, 2001	\$ 176,260
	<u> </u>
For the year ended December 31, 2002	\$ 52,429
	<u> </u>
For the year ended December 31, 2003	\$ 34,217
	<u> </u>

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Table of Contents**Estimated Amortization Expense**

For the year ended December 31, 2004	\$30,742
For the year ended December 31, 2005	\$27,703
For the year ended December 31, 2006	\$21,004
For the year ended December 31, 2007	\$15,138
For the year ended December 31, 2008	\$ 5,846

Goodwill

Increases in the carrying amount of goodwill are a result of acquisitions totaling \$60,738 (See Note 5). The segment detail is as follows:

	North America	International	Total
Balance as of January 1, 2002	2,899,398	206,324	3,105,722
Goodwill acquired, net	40,928	40,466	81,394
Currency Translation		5,535	5,535
Balance as of December 31, 2002	\$2,940,326	\$252,325	\$3,192,651
Goodwill acquired, net	24,925	35,813	60,738
Reclassifications	(14,398)	(865)	(15,263)
Currency Translation		50,222	50,222
Balance as of December 31, 2003	\$2,950,853	\$337,495	\$3,288,348

Had the Company determined amortization expense under SFAS No. 142 in 2001, the Company's net income in the year ended December 31, 2001 would have increased by \$102,775 (\$94,958 due to goodwill amortization, \$7,817 and \$7,817 due to amortization originally recorded on indefinite-life intangible assets) to \$166,129. All income per Ordinary and Preference share, as reported in December 31, 2001 would have increased by \$1.07 (\$0.99 due to goodwill amortization, \$0.08 due to amortization originally recorded on indefinite-life intangible assets) had the company adopted SFAS No. 142 as of January 1, 2001.

10. Accrued Expenses and Other Current Liabilities

As at December 31, 2003 and 2002 accrued expenses and other current liabilities consisted of the following:

	2003	2002
Accrued salaries and wages	\$143,747	\$121,212
Unapplied cash and receivable credits	65,624	63,773
Derivatives	51,446	28,656
Accrued insurance	45,015	48,165
Accrued operating expenses	41,236	33,369

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Accrued interest	39,448	35,861
Withholding tax and VAT	25,818	20,538
Accrued physician compensation	19,844	19,211
Commissions	17,568	14,877
Deferred income	10,336	8,359
Bonuses and Rebates	10,122	10,324
Accrued legal and compliance costs	7,767	5,821
Other	75,859	59,062
	<u> </u>	<u> </u>
Total accrued expenses and other current liabilities	\$553,830	\$469,228
	<u> </u>	<u> </u>

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Table of Contents**11. Debt and Capital Lease Obligations****Short term borrowings**

Short-term borrowings of \$89,417 and \$124,964 at December 31, 2003, and 2002, respectively, represent amounts borrowed by certain of the Company's subsidiaries under lines of credit with commercial banks. The average interest rates on these borrowings at December 31, 2003, and 2002 was 3.38% and 4.67%, respectively. For information regarding short-term borrowings from affiliates see Note 4b).

Excluding amounts available under the Senior Credit Agreement (as described below), at December 31, 2003, FMS had \$96,399 available under such commercial bank agreements. Some of these lines of credit are secured by the accounts receivable of FMS subsidiary that is party to the agreement and contain various covenants including, but not limited to, requirements for maintaining defined levels of working capital, net worth, capital expenditures and certain financial ratios.

Long-term borrowings

As of December 31, 2003 and 2002, long-term debt and capital lease obligations consisted of the following:

	<u>2003</u>	<u>2002</u>
Senior Credit Agreement	\$ 912,300	\$ 861,900
Capital leases	9,919	10,645
Euro Notes	162,296	134,758
Other	117,474	104,301
	<u>1,201,989</u>	<u>1,111,604</u>
Less current maturities	(90,365)	(22,394)
	<u>\$ 1,111,624</u>	<u>\$ 1,089,210</u>

2003 Senior Credit Agreement

On February 21, 2003, the Company entered into an amended and restated bank agreement (hereafter, the 2003 Senior Credit Agreement) with Bank of America N.A., Credit Suisse First Boston, Dresdner Bank AG New York, JPMorgan Chase Bank, The Bank of Nova Scotia and certain other lenders (collectively, the Lenders), replacing the 1996 Senior Credit Agreement that was scheduled to expire at September 30, 2003. Under the terms of the 2003 Senior Credit Agreement, the Lenders made available to the Company and certain subsidiaries and affiliates an aggregate amount of up to \$1,500,000. On August 22, 2003, the 2003 Senior Credit Agreement was amended (Amendment 1) so that, in effect, the aggregate amount of \$1,500,000 was voluntarily reduced to \$1,400,000 and the interest rate on a new term loan facility (Loan C, see below) was 25 basis points lower than on Loan B, which was repaid. The revolving loan facility and Loan A under the 2003 Senior Credit Agreement remain outstanding and were not affected by the amendment.

The credit facilities are:

a revolving credit facility of up to \$500,000 (of which up to \$250,000 is available for letters of credit, up to \$300,000 is available for borrowings in certain non-U.S. currencies, up to \$75,000 is available as swing lines in U.S. dollars, up to \$250,000 is available as a competitive loan facility and up to \$50,000 is available as swing lines in certain non-U.S. currencies, the total of which cannot exceed \$500,000) which will be due and payable on October 31, 2007.

a term loan facility (Loan A) of \$500,000, also scheduled to expire on October 31, 2007. The terms of the 2003 Senior Credit Agreement require payments that permanently reduce the term loan facility. The repayment begins in the third quarter of 2004 and amounts to \$25,000 per quarter. The remaining amount outstanding is due on October 31, 2007.

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a term loan facility (Loan B) of \$500,000 scheduled to expire in February 2010. Loan B was repaid as agreed in Amendment 1 to the 2003 Senior Credit Agreement under which the Lenders have made

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available to the Company a term loan facility (Loan C) in the amount of \$400,000. The proceeds of Loan C, together with cash on hand, were used to permanently repay Loan B under the 2003 Senior Credit Agreement.

a term loan facility (Loan C) of \$400,000 scheduled to expire February 21, 2010 subject to an early repayment requirement on October 31, 2007 if the Trust Preferred Securities due February 1, 2008 are not repaid or refinanced or their maturity is not extended prior to that date. The terms of Loan C require quarterly payments totaling \$1,000 per quarter beginning with the third quarter of 2003.

For the revolving credit facility and Loan A, interest is at a rate equal to LIBOR plus an applicable margin, or base rate, defined as the higher of the Bank of America prime rate or the Federal Funds rate plus 0.5% plus the applicable margin. The applicable margin is variable and depends on the ratio of the Company's funded debt to EBITDA as defined in the 2003 Senior Credit Agreement. The initial interest rate for Loan B was LIBOR plus 2.5%. Fees are also payable at a percentage (initially 0.50%) per annum on the portion of the revolving credit facility not used. The initial interest rate for Loan C is LIBOR plus 2.25% or the base rate plus 1.25%, which is 25 basis points less than the former Loan B. In addition to scheduled principal payments, indebtedness outstanding under the 2003 Senior Credit Agreement will be reduced by portions of the net cash proceeds from certain sales of assets, securitization transactions other than the Company's existing accounts receivable financing facility and the issuance of subordinated debt.

The 2003 Senior Credit Agreement contains affirmative and negative covenants with respect to the Company and its subsidiaries and other payment restrictions. Some of the covenants limit indebtedness of the Company and investments by the Company, and require the Company to maintain certain ratios defined in the agreement. Additionally, the 2003 Senior Credit Agreement provides for a dividend restriction which is \$150,000 for dividends paid in 2004, and increases in subsequent years. In default, the outstanding balance under the 2003 Senior Credit Facility becomes immediately due and payable at the option of the Lenders. As of December 31, 2003, the Company is in compliance with all financial covenants under the 2003 Senior Credit Agreement.

Euro Notes

In 2001, the Company issued four tranches of senior notes (Euro Notes) totaling 128,500 in aggregate principal amount. The first tranche was for 80,000 with a fixed interest rate of 6.16% and the second and third tranches were for 28,500 and 15,000, respectively, with variable interest rates that averaged 3.84.% in 2003 and 4.78% in 2002. The final tranche was for 5,000 at a fixed rate of 5.33%. All four tranches have a maturity date of July 13, 2005. Both floating rates are tied to the EURIBOR rate.

Annual Payments

Aggregate annual payments applicable to the 2003 Senior Credit Agreement, Euro Notes, capital leases and other borrowings for the five years subsequent to December 31, 2003 (excluding the Company's trust preferred securities) are:

2004	90,365
2005	293,017
2006	122,658
2007	282,813
2008	15,278
Thereafter	397,858
	\$ 1,201,989
	\$ 1,201,989

Table of Contents**12. Employee Benefit Plans****Defined Benefit Pension Plans**

The Company currently has two principal pension plans, one for German employees, the other covering employees in the United States. Plan benefits are generally based on years of service and final salary. Consistent with predominant practice in the Federal Republic of Germany, FMS's pension obligations in Germany are unfunded. In the United States NMC's non-contributory, defined benefit pension plan was curtailed in the first quarter of 2002. Each year FMCH contributes at least the minimum required by the Employee Retirement Income Security Act of 1974, as amended. There is no minimum funding requirement for FMCH for the defined benefit plan in 2004. The following tables provides a reconciliation of benefit obligations, plan assets, and funded status of the plans. Benefits paid as shown in the reconciliation of plan assets include only benefit payments from the Company's funded benefit plans.

	2003	2002	2001
Change in benefit obligation:			
Benefit obligation at beginning of year	\$ 184,468	\$ 169,623	\$ 145,308
Translation loss (gain)	8,870	6,484	(1,483)
Service cost	3,486	5,137	13,251
Interest cost	13,419	11,208	10,210
Curtailement	0	(22,216)	
Transfer of plan participants	1,356	84	(34)
Actuarial loss	33,563	17,764	6,127
Benefits paid	(3,922)	(3,616)	(3,756)
	<u>\$ 241,240</u>	<u>\$ 184,468</u>	<u>\$ 169,623</u>
Change on plan assets:			
Fair value of plan assets at beginning of year	\$ 83,191	\$ 89,845	\$ 88,817
Actual return on plan assets	13,898	(9,799)	(4,838)
Employer contributions	41,481	6,313	9,282
Benefits paid	(3,323)	(3,168)	(3,416)
	<u>\$ 135,247</u>	<u>\$ 83,191</u>	<u>\$ 89,845</u>
Funded status:			
Unrecognized net loss	\$ 105,994	\$ 101,277	\$ 79,778
Unrecognized prior service cost	(61,595)	(37,302)	(10,232)
Unrecognized transition obligation		(85)	(143)
	<u>\$ 44,399</u>	<u>\$ 63,890</u>	<u>\$ 69,406</u>
Calculation of Additional Minimum Liability⁽¹⁾			
Fair Value of plan assets	\$ 135,247	\$ 83,191	\$ 89,845
Accumulated benefit obligation (ABO)	184,489	142,893	137,953
	<u>\$ 49,242</u>	<u>\$ 59,702</u>	<u>\$ 48,108</u>

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	2003	2002	2001
Accrued benefit costs	\$ (6,411)	\$ 27,440	\$42,639
Additional Minimum Liability	\$ 55,653	\$ 32,262	\$ 5,469
Thereof accumulated other comprehensive income	\$ 55,653	\$ 32,262	\$ 5,469
Total pension liability (at December 31)	\$100,052	\$ 96,152	\$74,875
Weighted average assumptions for benefit obligation as of December 31:			
Discount rate	6.14%	6.53%	7.22%
Rate of compensation increase	4.27%	4.28%	4.50%
Components of net period benefit cost:			
Service cost	\$ 3,486	\$ 5,137	\$13,251
Interest cost	13,419	11,208	10,210
Expected return on plan assets	(7,688)	(8,102)	(8,815)
Amortization of transition obligation	92	77	73
Amortization unrealized losses	3,971	183	188
Curtailment gain		(12,620)	
Net amortization			(1,377)
Net periodic benefit costs	\$ 13,280	\$ (4,117)	\$13,530
Weighted average assumptions for net periodic benefit cost for the year ended December 31			
Discount rate	6.52%	7.12%	7.22%
Expected return of plan assets	8.50%	9.00%	9.94%
Rate of compensation increase	4.27%	4.28%	4.50%

(1) this calculation refers only to companies with ABO in excess of plan assets

Plan Investment Policy and Strategy

The investment strategy for the FMS North America pension plan is to earn a long-term rate of return on assets of at least 7.5% compounded annually while utilizing a target investment allocation of 50% equities and 50% long-term U.S. bonds.

The investment policy considers that there will be a time horizon for invested funds of more than 5 years. The total portfolio will be measured against a policy index that reflects the asset class benchmarks and the target asset allocation. The performance benchmarks for the separate asset classes include: S&P 500 Index, Russell 2000 Growth Index, MSCI EAFE Index, Lehman U.S. Long Government/ Credit bond Index and the HFRI Fund of Funds Index. The following schedule describes FMCH's allocation for its plans:

	Allocation 2003 in %	Allocation 2002 in %	Target allocation in %
Categories of plan assets			
Equity securities	52%	56%	50%
Debt securities	48%	44%	50%
Total	100%	100%	100%

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	Long term expected return
Overall expected long-term rate	7.50%
	2004
Expected total contributions to plan assets for following year	\$ 10,702

The measurement date used to determine pension benefit measurements was December 31, 2003 for the plans in the United States and September 30, 2003 for the non-U.S. plans.

Defined Contribution Plans

FMCH's employees are eligible to join 401(k) savings plan. The Company's total contributions for the years ended December 31, 2003, 2002 and 2001 was \$14,754, \$12,974 and \$10,647, respectively.

13. Mandatorily Redeemable Trust Preferred Securities

The Company originally issued Trust Preferred Securities through five Fresenius Medical Care Capital Trusts, statutory business trusts organized under the laws of the State of Delaware. FMS owns all of the common securities of these trusts. The sole asset of each trust is a senior subordinated note of a wholly-owned subsidiary of FMS and related guarantees by FMS, Fresenius Medical Care Deutschland GmbH (D-GmbH) and FMCH; D-GmbH and FMCH being the Guarantor Subsidiaries. The Trust Preferred Securities are guaranteed by FMS through a series of undertakings by the Company and the Subsidiary Guarantors.

The Trust Preferred Securities entitle the holders to distributions at a fixed annual rate of the stated amount and are mandatorily redeemable after 10 years. Earlier redemption may also occur upon a change of control followed by a rating decline or defined events of default including a failure to pay interest. Upon liquidation of the trusts, the holders of Trust Preferred Securities are entitled to a distribution equal to the stated amount. The Trust Preferred Securities do not hold voting rights in the trust except under limited circumstances.

On February 14, 2002, the Company redeemed the entire \$360,000 aggregate liquidation amount outstanding of its 9% Trust Preferred Securities due 2006. The terms of the securities, which were issued in 1996, provided for optional redemption commencing December 1, 2001 at a redemption price of 104.5% of the liquidation amount, plus distributions accrued to the redemption date. The Company redeemed the securities at a price of \$1,045 per \$1,000 liquidation amount plus accrued distributions of \$18.25 per \$1,000. At that time an extraordinary loss of \$11,777 was recorded as a result of the early redemption of debt, consisting of \$16,200 of redemption premium and \$3,317 of write-off of associated debt issuance costs, net of a \$7,740 tax benefit.

As of January 1, 2003 the Company adopted Statement of Financial Accounting Standards (SFAS) No. 145, *Rescission of FASB Statements No. 4, 44, and 64, Amendment of FASB Statement No. 13, and Technical Corrections* in regard to SFAS No. 4. As a result, the loss on the early redemption of the 9% Trust Preferred Securities is no longer presented as an extraordinary loss, but is presented in interest expense, with the related income tax effect included in income taxes.

The Trust Preferred Securities outstanding as of December 31 are as follows:

	Year Issued	Stated Amount	Interest Rate	Mandatory Redemption Date	2003	2002
Fresenius Medical Care Capital Trust II	1998	\$450,000	7 7/8%	February 1, 2008	450,000	450,000
Fresenius Medical Care Capital Trust III	1998	DM 300,000	7 3/8%	February 1, 2008	193,728	160,858
Fresenius Medical Care Capital Trust IV	2001	\$225,000	7 7/8%	June 15, 2011	222,150	221,766

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Fresenius Medical Care Capital Trust V	2001	300,000	7 3/8%	June 15, 2011	<u>376,439</u>	<u>312,657</u>
					<u>\$1,242,317</u>	<u>\$1,145,281</u>

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Table of Contents**14. Minority Interests**

At December 31, 2003 and 2002, minority interests were as follows:

	<u>2003</u>	<u>2002</u>
FMCH Preferred Stock:		
Preferred Stock, \$100 par value		
6% Cumulative; 40,000 shares authorized; 36,460 issued and outstanding	\$ 3,646	\$ 3,646
8% Cumulative Class A; 50,000 shares authorized; 16,176 issued and outstanding	1,618	1,618
8% Noncumulative Class B; 40,000 shares authorized; 21,483 issued and outstanding	2,148	2,148
Preferred Stock, \$0.10 par value		
Noncumulative Class D; 100,000,000 shares authorized; 89,062,316 issued and outstanding		8,906
	<u>7,412</u>	<u>16,318</u>
Sub-total FMCH minority interest	7,412	16,318
Other minority interest	6,693	6,204
	<u>6,693</u>	<u>6,204</u>
Total minority interest	<u>\$ 14,105</u>	<u>\$ 22,522</u>

On February 4, 2003, the Company and FMCH announced FMCH was exercising its right to redeem all of the outstanding shares of the Class D Preferred Stock (Class D Shares) of FMCH. The Class D Shares were issued to the common shareholders of W.R. Grace & Co. in connection with the 1996 combination of the worldwide dialysis business of Fresenius AG with the dialysis business of W.R. Grace to form the Company.

Commencing on March 28, 2003, Class D Shares that were properly transferred to and received by the redemption agent were redeemed at a redemption price of \$0.10 per share. FMCH redeemed the 89 million outstanding Class D Shares at a total cash outflow of approximately \$8,900. This transaction had no earnings impact for the Company. After March 28, 2003 the Class D Shares ceased to be issued and outstanding shares of FMCH's capital stock.

15. Shareholders Equity**Capital Stock**

As of December 31, 2003, the Company's capital stock consisted of 26,213,979 Preference shares (53,597,700 shares authorized) without par value with a nominal amount of 2.56 per share totaling \$69,616 and of 70,000,000 Ordinary shares without par value with a nominal amount of 2.56 per share totaling \$229,494.

As of December 31, 2002 and 2001, the Company's capital stock was divided into 26,188,575 and 26,176,508 Preference shares (53,597,700 shares authorized) amounting to \$69,540 and \$69,512 respectively and 70,000,000 Ordinary shares amounting to \$229,494.

Under the German Stock Corporation Act, the shareholders of a stock corporation may empower the management board to issue shares in a specified aggregate nominal value not exceeding 50% of the issued share capital at the time of the passing of the resolution, in the form of Conditional Capital (*bedingtes Kapital*) or Approved Capital (*genehmigtes Kapital*). The authorization for the issuance of Approved Capital is limited for a period not exceeding five years from the date the shareholders' resolution becomes effective.

The authorized and issued number of Preference shares was impacted during the fiscal years 2003, 2002 and 2001 by the following transactions:

Approved Capital

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By resolution of the annual general meetings on May 30, 2000 and May 23, 2001, respectively, the management board, with the approval of the supervisory board, was authorized to increase nominal share capital by the maximum amount of:

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30,720, corresponding to 12,000,000 Preference shares, by issuing new non-voting Preference shares for cash, new Approved Capital I. As of December 31, 2003, 12,000,000 Preference shares are available for issuance under Approved Capital I.

20,480, corresponding to 8,000,000 Preference shares, by issuing new non-voting Preference shares for cash or against contributions in kind, new Approved Capital II. As of December 31, 2003, 8,000,000 Preference shares are available for issuance under Approved Capital II.

The authorizations of Approved Capital I and Approved Capital II are effective until May 29, 2005 and May 22, 2006, respectively.

The management board may exclude statutory preemptive rights in connection with the issuance of Preference shares using Approved Capital II if the shares are issued against a contribution in kind to acquire a company or an interest in a company or if the shares are issued for cash and the issue price is not materially lower than the price of such shares on the stock exchange.

Conditional Capital

By resolution of the general meeting on May 23, 2001, FMS's share capital was conditionally increased by up to 10,240, divided into a maximum of 4,000,000 new non-voting Preference shares. This conditional capital increase may be issued only upon exercise of grants by employees under the FMC 2001 International Stock Incentive Plan. As of December 31, 2003 all 4,000,000 Preference shares are available for issue.

In addition, conditional capital of a nominal amount of up to 9,216 representing 514,505 non-voting Preference shares is available for employees exercising rights granted under other stock-based compensation plans.

Dividends

Under the German Stock Corporation Act, the amount of dividends available for distribution to shareholders is based upon the unconsolidated retained earnings of Fresenius Medical Care AG as reported in its balance sheet determined in accordance with the German Commercial Code (*Handelsgesetzbuch*).

If no dividends were declared for two consecutive years after the year for which the Preference shares are entitled to dividends, then the holders of such Preference shares would be entitled to the same voting rights as holders of Ordinary shares until all arrearages are paid. In addition, the payment of dividends by FMS is subject to limitations under the 2003 Senior Credit Agreement (see Note 11).

Cash dividends of \$107,761 for 2002 in the amount of 1.00 per Preference share and 0.94 per Ordinary share were paid on May 23, 2003.

Cash dividends of \$76,743 for 2001 in the amount of 0.91 per Preference share and 0.85 per Ordinary share were paid on May 23, 2002.

Cash dividends of \$65,782 for 2000 in the amount of 0.84 per Preference share and 0.78 per Ordinary share were paid on May 24, 2001.

Table of Contents**16. Earnings Per Share**

The following table is a reconciliation of the numerators and denominators of the basic and diluted earnings per share computations. At December 31, 2001, the performance criteria for the 1998 Plan stock options granted in 2000 had not been met. Due to this, the stock options granted are excluded from the diluted earnings per share computations.

	<u>2003</u>	<u>2002</u>	<u>2001</u>
<i>Numerators:</i>			
Net income	\$ 331,180	\$ 289,790	\$ 63,354
less:			
Preference on Preference shares	1,778	1,485	1,399
Income available to all class of shares	<u>\$ 329,402</u>	<u>\$ 288,305</u>	<u>\$ 61,955</u>
<i>Denominators:</i>			
Weighted average number of:			
Ordinary shares outstanding	70,000,000	70,000,000	70,000,000
Preference shares outstanding	26,191,011	26,185,178	26,035,330
Total weighted average shares outstanding	96,191,011	96,185,178	96,035,330
Potentially dilutive Preference shares	145,861	66,120	399,697
Total weighted average shares outstanding assuming dilution	96,336,872	96,251,298	96,435,027
Total weighted average Preference shares outstanding assuming dilution	26,336,872	26,251,298	26,435,027
Basic income per Ordinary share	\$ 3.42	\$ 3.00	\$ 0.65
Plus preference per Preference share	0.07	0.06	0.05
Basic income per Preference Share	<u>\$ 3.49</u>	<u>\$ 3.06</u>	<u>\$ 0.70</u>
Fully diluted income per Ordinary share	\$ 3.42	\$ 3.00	\$ 0.64
Plus preference per Preference share assuming dilution	0.07	0.06	0.05
Fully diluted income per Preference share	<u>\$ 3.49</u>	<u>\$ 3.06</u>	<u>\$ 0.69</u>

17. Stock Options

In connection with the formation of FMS in 1996, certain options outstanding under stock option plans of W.R. Grace and FUSA were exchanged, for equivalent options with respect to FMS Ordinary shares (the FMC Rollover Plan).

During the year ended December 31, 2003, 79,491 FMC Rollover Plan options were exercised by employees. In connection therewith, Fresenius AG transferred 26,497 Ordinary shares to employees and remitted \$631 to the Company. The \$631 has been accounted for as a capital contribution within additional paid in capital. Rollover Plan options for 24,927 Ordinary American Depository Shares were exercisable as of December 31, 2003 at a weighted average exercise price of \$13.54.

Fresenius Medical Care Plan

In connection with the formation of the Company, FMS adopted a stock incentive plan (the FMC Plan) for FMS's key management and executive employees. The options have a ten year term and vest after three or five years. During 2003, no options were exercised. As of December 31, 2003, 53,389 options for preference shares were exercisable with a price range between \$55.59 and \$78.33 per share. Effective

September 2001, no additional awards could be granted under the FMC Plan.

Table of Contents***Fresenius Medical Care 98 Plan 1 and Plan 2***

During 1998, the Company adopted two stock incentive plans (FMC 98 Plan 1 and FMC 98 Plan 2) for FMS's key management and executive employees. Under FMC 98 Plan 1, eligible employees have the right to acquire Preference shares of the Company. Options granted under FMC 98 Plan 1 have a ten year term, and one third of them vest on each of the second, third and fourth anniversaries of the award date. The maximum number of Preference shares that may be issued under this plan is 2,443,333 less any shares issued under the FMC Plan. Any shares available due to forfeiture of grants under the FMC Plan would be considered available under FMC 98 Plan 1 as long as the total Preference shares issued under both plans does not exceed the 2,443,333 shares noted above.

Under FMC 98 Plan 2, eligible employees have the right to acquire Preference shares (the Options) of the Company. The share price of the Preference share shall be equal to the average of the official daily quotation prices of the Preference shares on the Frankfurt Stock Exchange on the thirty days (30) of trading immediately prior to the date of grant of the Option. One third of an Option vests on each of the second, third and fourth anniversaries of the award date, provided that the Company achieves certain performance criteria for the full fiscal year following the grant date in comparison to its performance for the full fiscal year preceding the grant date. Options granted under FMC 98 Plan 2 have a 10-year term. The maximum number of Preference shares that may be issued under this plan is 2,500,000 shares, of which 500,000 are designated for management board members and 2,000,000 are for other managerial staff. Each Option is exercisable for one Preference share.

The following table shows the number of Preference shares available and the price range (in \$ and) under FMC 98 Plan 1 and FMC 98 Plan 2:

	Options (in thousands)	Average price range	Average price range
<i>FMC 98 Plan 1</i>			
Balance at December 31, 2001	1,690	32.90-56.24	\$41.55-71.03
Granted			
Exercised	10	32.90-40.70	41.55-51.40
Forfeited	65	32.90-56.24	41.55-71.03
Balance at December 31, 2002	1,615	32.90-56.24	41.55-71.03
Exercised	8	32.90	41.55
Forfeited	110	32.90-56.24	41.55-71.03
Balance at December 31, 2003	1,497	32.90-56.24	\$41.55-71.03
Exercisable at December 31, 2003	1,446	32.90-56.24	\$41.55-71.03
<i>FMC 98 Plan 2</i>			
Balance at December 31, 2001	782	32.41-47.64	\$40.93-60.17
Granted			
Exercised	2	32.41-44.66	40.93-56.41
Forfeited	301	32.41-47.64	40.93-60.17
Balance at December 31, 2002	479	32.41-44.66	40.93-56.41
Exercised	17	32.41	40.93
Forfeited	26	32.41-44.66	40.93-56.41
Balance at December 31, 2003	436	32.41-44.66	\$40.93-56.41

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Exercisable at December 31, 2003	<u>436</u>	<u>32.41-44.66</u>	<u>\$40.93-56.41</u>
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The following table summarizes information about stock options outstanding for both 98 Plans at December 31, 2003:

Range of exercise prices in \$	Options outstanding	Weighted average remaining contractual life	Weighted average exercise price	Options exercisable	Weighted average exercise price
\$40.01-45.00	565,038	5.50	\$41.29	565,038	\$41.29
\$45.01-50.00					
\$50.01-55.00	117,217	5.90	51.63	117,217	51.63
\$55.01-60.00	197,618	4.50	56.41	197,618	56.41
\$60.01-65.00	531,438	6.63	61.83	486,117	61.84
\$65.01-70.00	16,712	7.25	66.05	11,142	66.05
\$70.01-75.00	504,771	4.40	71.03	504,771	71.03
	1,932,794	5.46	\$57.09	1,881,903	\$56.95

Proceeds totaling \$969 from exercise of 8,648 shares under FMC 98 Plan 1 and 16,756 shares under FMC 98 Plan 2 in 2002 were recorded as a capital contribution. Effective September 2001, no additional grants or options can be awarded under FMC 98 Plan 1 or FMC 98 Plan 2.

Fresenius Medical Care 2001 International Stock Incentive Plan

On May 23, 2001, by resolution of the annual general meeting, the FMC 98 Plans were replaced by a new plan. Under the terms of this new plan, convertible bonds with a principal of up to 10,240 may be issued to the members of the management board and other employees of the Company representing grants for up to 4 million non-voting Preference shares. The convertible bonds have a par value of 2.56 and are interest bearing at a rate of 5.5%. Purchase of the bonds may be funded by a non-recourse loan secured by the bond with respect to which the loan was made. The Company has the right to offset its obligation on a convertible bond against the employee obligation on the related loan; therefore, the convertible bond obligations and employee loan receivables are not reflected in the Company's consolidated financial statements. The bonds mature in ten years and are generally convertible after four years. The bonds may be issued either as convertible bonds which are subject to a stock price target or convertible bonds without a stock price target. In the case of convertible bonds which are subject to a stock price target the conversion right is only exercisable if the quoted price of the Preference shares exceeds the quoted price at grant date by at least 25% at any given date subsequent to the date of the grant. Participants have the right to choose between convertible bonds with or without the stock price target. The number of convertible bonds awarded to those employees who select the bonds without a stock price target will be reduced by 15%. Each convertible bond entitles the holder thereof, upon payment of a conversion price to convert the bond into one Preference share. The conversion price of the convertible bonds which are not subject to the stock price target is determined by the average price of the Preference shares during the last 30 trading days prior to the date of grant. Up to 20% of the total amount available for the issuance of convertible bonds may be issued each year through May 22, 2006.

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The following table presents the number of Preference shares available and the average price range (in \$ and) under the FMC 2001 International Stock Incentive Plan.

	Bonds (in thousands)	Average price range	Average price range
<i>FMC International Plan</i>			
Balance at December 31, 2001	720	53.27-73.72	67.28-93.11
Granted	771	25.13-43.16	31.74-54.51
Forfeited	23	56.42-73.72	71.26-93.11
Balance at December 31, 2002	1,468	25.13-73.72	\$31.74-93.11
Granted	622	29.10-37.02	36.75-46.76
Forfeited	87	25.13-73.72	31.74-93.11
Balance at December 31, 2003	2,003	25.13-73.72	\$31.74-93.11
Exercisable at December 31, 2003	212	53.27-73.72	\$67.28-93.11

Fair Value of Stock Options

The per share weighted-average fair value of stock options granted during 2003, 2002 and 2001 was \$14.26, \$11.11 and \$16.76, respectively, on the date of the grant using the Black-Scholes option-pricing model with the weighted-average assumptions presented below.

	2003	2002	2001
Weighted-average assumptions:			
Expected dividend yield	2.60%	2.20%	1.50%
Risk-free interest rate	3.80%	3.80%	4.90%
Expected volatility	40.00%	40.00%	40.00%
Expected life of options	5.3 years	5.3 years	5.3 years

The Company applies APB Opinion No. 25 in accounting for stock compensation and, accordingly, recognized compensation expense of \$1,456, \$1,126 and \$1,153 for stock options granted in 2003, 2002 and 2001.

18. Income Taxes

Income before income taxes and minority interest is attributable to the following geographic locations:

	2003	2002	2001
Germany	\$ 78,124	\$ 86,701	\$ 123,141
United States	368,382	289,954	(60,930)
Other	99,170	91,795	94,077
	\$545,676	\$468,450	\$156,288

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Income tax expense (benefit) for the years ended December 31, 2003, 2002, and 2001, consisted of the following:

	<u>2003</u>	<u>2002</u>	<u>2001</u>
Current:			
Germany	\$ 51,849	\$ 29,367	\$ 30,094
United States	22,346	53,878	87,923
Other	35,505	32,124	31,079
	<u>109,700</u>	<u>115,369</u>	<u>149,097</u>
Deferred:			
Germany	(1,280)	10,069	7,651
United States	102,142	47,437	(72,455)
Other	2,152	2,198	6,909
	<u>103,014</u>	<u>59,705</u>	<u>(57,895)</u>
	<u>\$ 212,714</u>	<u>\$ 175,074</u>	<u>\$ 91,202</u>

Under the provisions applicable as a result of the Flood Victim Solidarity Law, for the fiscal year ended December 31, 2003 the Company is subject to German federal corporation income tax at a base rate of 26.5% (25% in 2002) plus a solidarity surcharge of 5.5% on federal corporation taxes payable. Because of this, the statutory rate for the year ended December 31, 2003 amounted to 27.96% compared to 26.375% in 2002.

The increase of the base rate of German federal corporation income tax from 25% to 26.5% was enacted by the German government with the Flood Victim Solidarity Law in September 2002. This increase was effective only for 2003 and the tax rate returned to 25% on January 1, 2004.

The difference in income tax expense from the amounts computed by applying the German federal corporation income tax rate, including the solidarity surcharge, on income before income taxes and minority interest (27.96% for fiscal year 2003, 26.375% for fiscal years 2002 and 2001, respectively) is as follows:

	<u>2003</u>	<u>2002</u>	<u>2001</u>
Computed expected income tax expense at the undistributed earnings rate	\$ 152,571	\$ 123,554	\$ 41,221
Trade income taxes, net of German federal corporation income tax benefit	15,486	12,184	13,663
U.S. State income taxes, net of federal tax benefit	13,535	10,740	420
Tax free income	(12,155)	(11,078)	(5,327)
Non-deductible portion of special charge for legal matters			14,216
Amortization of non-tax deductible goodwill			19,678
Foreign tax rate differential	29,904	25,929	7,538
Non-deductible expenses	6,993	7,827	691
Other	6,380	5,918	(898)
	<u>212,714</u>	<u>175,074</u>	<u>91,202</u>
Provision for income taxes	<u>\$ 212,714</u>	<u>\$ 175,074</u>	<u>\$ 91,202</u>
Effective tax rate	<u>39.0%</u>	<u>37.4%</u>	<u>58.4%</u>

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The tax effects of the temporary differences that give rise to deferred tax assets and liabilities at December 31 are presented below:

	<u>2003</u>	<u>2002</u>
Deferred tax assets:		
Accounts receivable, primarily due to allowance for doubtful accounts	\$ 30,939	\$ 25,962
Inventory, primarily due to additional costs capitalized for tax purposes, and inventory reserve accounts	28,126	21,368
Accrued expenses and other liabilities for financial accounting purposes, not currently tax deductible	110,921	129,154
Special charge for legal matters	48,199	46,580
Net operating loss carryforwards	40,237	47,971
Derivatives	27,685	42,370
Other	5,327	3,975
	<u> </u>	<u> </u>
Total deferred tax assets	\$ 291,432	\$ 317,380
Less: valuation allowance	(28,084)	(23,229)
	<u> </u>	<u> </u>
Net deferred tax assets	\$ 263,348	\$ 294,151
	<u> </u>	<u> </u>
Deferred tax liabilities:		
Accounts receivable, primarily due to allowance for doubtful accounts	\$ 32,003	\$ 20,207
Inventory, primarily due to inventory reserve accounts for tax purposes	8,706	6,646
Accrued expenses and other liabilities deductible for tax prior to financial accounting recognition	19,212	23,256
Plant and equipment, principally due to differences in depreciation	213,907	165,264
Derivatives	36,612	31,551
Other	14,251	9,006
	<u> </u>	<u> </u>
Total deferred tax liabilities	324,691	255,930
	<u> </u>	<u> </u>
Net deferred tax (liabilities) assets	\$ (61,343)	\$ 38,221
	<u> </u>	<u> </u>

During 2003, the valuation allowance increased by \$4,856. In 2002, the valuation allowance increased by \$16,800 mainly attributable to currency exchange losses in Latin America.

The expiration of net operating losses is as follows:

2004	\$ 5,523
2005	12,165
2006	4,712
2007	9,043
2008	10,528
2009	11,929
2010	5,950
2011	2,477
2012	4,152
2013	1,764
Thereafter	40,874
	<u> </u>
Total	\$ 109,117
	<u> </u>

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In assessing the realizability of deferred tax assets, management considers whether it is more likely than not that some portion or all of the deferred tax assets will not be realized. The ultimate realization of deferred tax assets is dependent upon the generation of future taxable income during the periods in which those temporary differences become deductible. Management considers the scheduled reversal of deferred tax liabilities and projected future taxable income in making this assessment. Based upon the level of historical taxable income and projections for future taxable income over the periods in which the deferred tax assets are deductible, management believes it is more likely than not the Company will realize the benefits of these deductible differences, net of the existing valuation allowances at December 31, 2003.

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Provision has not been made for additional taxes on approximately \$267,763 undistributed earnings of foreign subsidiaries. The majority of these earnings have been, and will continue to be, permanently reinvested. The earnings could become subject to additional tax if remitted or deemed remitted as dividends; however, calculation of such additional tax is not practical. For fiscal years ending in 2004 and afterwards, dividends from German subsidiaries are 95% tax-exempt, i.e. 5% of dividend income is taxable for corporate tax purposes after recent German tax law changes. The effects of this new rule are estimated by management as negligible, as the majority of German investments are consolidated for tax purposes.

A 5% income inclusion has also been introduced on capital gains realized from the disposition of shares in German and foreign corporations and applies to fiscal years ending in 2004. Management does not anticipate significant additional income taxation.

19. Operating Leases

The Company leases buildings and machinery and equipment under various lease agreements expiring on dates through 2016. Rental expense recorded for operating leases for the years ended December 31, 2003, 2002 and 2001 was \$303,060, \$270,082, and \$237,174, respectively.

In December 2003, the Company exercised an option to terminate an operating lease for certain manufacturing equipment in its Ogden, Utah, North American facility. The equipment was purchased for approximately \$66,000 and is reflected as a capital expenditure in the accompanying consolidated statement of cash flows.

Future minimum rental payments under noncancelable operating leases for the five years succeeding December 31, 2003 and thereafter are:

2004	\$ 237,239
2005	208,087
2006	172,876
2007	119,575
2008	88,377
Thereafter	265,117
	<hr/>
	\$ 1,091,271
	<hr/>

20. Legal Proceedings***Commercial Litigation***

The Company was formed as a result of a series of transactions pursuant to the Agreement and Plan of Reorganization (the Merger) dated as of February 4, 1996 by and between W.R. Grace & Co. and Fresenius AG. At the time of the Merger, a W.R. Grace & Co. subsidiary known as W.R. Grace & Co.-Conn. had, and continues to have, significant potential liabilities arising out of product-liability related litigation, pre-Me