Interim financial report (US-GAAP) 2nd quarter 2016

Fresenius Medical Care AG & Co. KGaA

Hof an der Saale

Germany



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FINANCIAL INFORMATION

Management's Discussion and Analysis

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 (the "Exchange Act"). When used in this report, the words "outlook," "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, and 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. We have based these forward-looking statements on current estimates and assumptions made to the best of our knowledge. By their nature, such forward-looking statements involve risks, uncertainties, assumptions and other factors which could cause actual results, including our financial condition and profitability, to differ materially positively or negatively relative to the results expressly or implicitly described in or suggested by these statements. Moreover, forward-looking estimates or predictions derived from third parties' studies or information may prove to be inaccurate. Consequently, we cannot give any assurance regarding the future accuracy of the opinions set forth in this report or the actual occurrence of the projected developments described herein. In addition, even if our future results meet the expectations expressed here, those results may not be indicative of our performance in future periods.

These risks, uncertainties, assumptions, and other factors that could cause actual results to differ from our projected results include, among others, the following:

- changes in governmental and commercial insurer reimbursement for our complete products and services portfolio, including the United States ("U.S.")
 Medicare reimbursement system for dialysis services;
- the outcome of government and internal investigations as well as litigation;
- risks relating to compliance with the government regulations applicable to our business including, in the U.S., the Anti-Kickback Statute, the False Claims Act, the Stark Law and the Foreign Corrupt Practices Act, the Food, Drug and Cosmetic Act and comparable regulatory regimes in many of the more than 120 countries in which we supply health care services and/or products;
- the influence of commercial insurers and managed care organizations, including
 efforts by these organizations to manage costs by limiting healthcare benefits,
 reducing provider reimbursement and/or restricting options for patient funding of
 premiums;
- the impact of health care reforms;
- product liability risks;
- risks relating our ability to continue to make acquisitions;
- the impact of currency fluctuations;
- changes in utilization patterns for pharmaceuticals and in our costs of purchasing pharmaceuticals;
- introduction of generic or new pharmaceuticals that compete with our pharmaceutical products;

- changes in raw material and energy costs or the ability to procure raw materials;
- collectability of our receivables, which depends primarily on the financial stability and liquidity of our governmental and commercial payors;
- our ability to achieve cost savings in various health care risk management programs in which we participate or intend to participate; and
- the greater size, market power and experience of certain competitors in certain geographic regions and business lines.

Important factors that could contribute to such differences are noted in "Financial Condition and Results of Operations – Overview, legislation and growth - Overview" below, in Note 10 in this report, in Note 19 of the Annual Report 2015 (Chapter 4) and in the section "Risk and Opportunities Report" in Chapter 2 of our Annual Report 2015.

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.

Our reported financial condition and results of operations are sensitive to accounting methods, assumptions and estimates that are the basis of our financial statements. The actual accounting policies, the judgments made in the selection 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 our financial statements and the discussion under "Financial Condition and Results of Operations - Results of Operations" below. There have been no significant changes during the six months ended June 30, 2016 to the items disclosed within the Annual Report 2015.

Financial Condition and Results of Operations

You should read the following discussion and analysis of the results of operations of Fresenius Medical Care AG & Co. KGaA ("FMC-AG & Co. KGaA," or the "Company") and its subsidiaries in conjunction with our unaudited consolidated financial statements and related notes contained elsewhere in this report and our disclosures and discussions in our Annual Report 2015. The results within this discussion and analysis are unaudited. In this report, "FMC-AG & Co. KGaA," or the "Company," "we," "us" or "our" refers to the Company or the Company and its subsidiaries on a consolidated basis, as the context requires. The term "North America Segment" refers to our North America operating segment; the term "EMEA Segment" refers to the Europe, Middle East and Africa operating segment, the term "Asia-Pacific Segment" refers to our Asia-Pacific operating segment, and the term "Latin America Segment" refers to our Latin America operating segment. The term "Corporate" includes certain headquarters' overhead charges, including accounting and finance, centrally managed production, asset management, quality management, procurement and research and development. The term "Constant Currency" or at "Constant Exchange Rates" means that we have translated local currency revenues for the current reporting period into U.S. dollars using the same average foreign currency exchange rates for the conversion of revenues into U.S. dollars that we used to translate local currency revenues for the comparable reporting period of the prior year, as described below under "Non - US GAAP Measures for Presentation - Constant Currency."

Overview, legislation and growth

Overview

We are the world's largest kidney dialysis company. We provide dialysis care and related services to persons who suffer from end stage renal disease ("ESRD") as well as

other health care services. We develop and manufacture a full range of dialysis machines, systems and disposable products, which we sell to customers in more than 120 countries and also use in our internal health care service operations. Our dialysis business is vertically integrated, providing dialysis treatment at our own dialysis clinics and supplying these clinics with a broad range of products. In addition, we sell dialysis products to other dialysis service providers. We describe our other health care services as "Care Coordination." Care Coordination currently includes coordinated delivery of pharmacy services, vascular, cardiovascular and endovascular specialty services, nondialysis laboratory testing services, physician services, hospitalist and intensivist services, health plan services and urgent care services, which, together with dialysis care services represent our health care services. Based on publicly reported sales and number of patients treated, our health care operations in dialysis services and dialysis products make us the world's largest kidney dialysis company. We estimated the volume of the global dialysis market was approximately \$73 billion in 2015. Dialysis patient growth results from factors such as the aging population and increased life expectancies; shortage of donor organs for kidney transplants; increasing incidence of kidney disease and better treatment of and survival of patients with diabetes, hypertension and other illnesses, which frequently lead to the onset of chronic kidney disease; improvements in treatment quality, which prolong patient life; and improving standards of living in developing countries, which make life-saving dialysis treatment available.

As a global company delivering health care services and dialysis products we face the challenge of addressing the needs of a wide variety of stakeholders, such as patients, customers, payors, regulators and legislators in very different economic environments and health care systems. In general, government-funded programs (in some countries in coordination with private insurers) pay for certain health care items and services provided to their citizens. Not all health care systems provide for dialysis treatment. Therefore, the reimbursement systems in various countries and ancillary services utilization environment significantly influence our business.

The majority of health care services we provide are paid for by governmental institutions. Approximately 32% of our consolidated revenues are attributable to U.S. federally-funded health care benefit programs, such as Medicare and Medicaid reimbursement, under which reimbursement rates are set by the Centers for Medicare & Medicaid Services ("CMS"). Legislative changes could affect Medicare reimbursement rates for a significant portion of the services we provide. To date, while we have generally experienced stable reimbursement globally, the stability of reimbursement in the U.S. has been affected by (i) the implementation of the ESRD prospective payment system ("ESRD PPS") in the U.S. in January 2011, (ii) the U.S. federal government across the board spending cuts in payments to Medicare providers commonly referred to as "U.S. Sequestration," (iii) the reduction to the ESRD PPS rate to account for the decline in utilization of certain drugs and biologicals associated with dialysis pursuant to the American Taxpayer Relief Act of 2012 ("ATRA") and (iv) the enactment of Protecting Access to Medicare Act of 2014 ("PAMA"). Please see the broader discussion of these legislative developments below:

Significant Legislative Impacts on U.S. Reimbursement

• Under Medicare Improvements for Patients and Providers Act of 2008 ("MIPPA"), for patients with Medicare coverage, all ESRD payments for dialysis treatments are made under a single bundled payment rate which provides a fixed payment rate, ESRD PPS, to encompass substantially all goods and services provided during the dialysis treatment. MIPPA further created the ESRD quality incentive program ("QIP") which dictates that dialysis facilities that fail to achieve quality standards established by CMS could have payments reduced by up to 2 percent.

- MIPPA also includes a provision for an annual adjustment to the ESRD PPS base rate based on changes in the costs of a "market basket" of certain health care items and services, less a productivity adjustment.
- Additionally, as a result of the Budget Control Act of 2011 ("BCA") and subsequent activity in Congress, a \$1.2 trillion sequester (across-the-board spending cuts) in discretionary programs took effect on March 1, 2013 and is expected to continue through mid-2024. In particular, a 2% reduction to Medicare payments took effect on April 1, 2013 and continues in force. Spending cuts pursuant to U.S. Sequestration have adversely affected and will continue to adversely affect our revenues, earnings and cash flows.
- In 2014, as mandated by ATRA, CMS issued a final rule for the ESRD PPS, which phased in payment reductions to account for changes in utilization of certain drugs and biologicals that are included in the ESRD PPS, which were subsequently modified by PAMA. These reductions will reduce our market basket inflation adjustment by 1.25% in 2016 and 2017, and 1% in 2018.

Significant Administrative Impacts on U.S. Reimbursement

On November 6, 2015, CMS published a final rule to update payment policies and rates under the ESRD PPS for renal dialysis services furnished on or after January 1, 2016. In this final rule, CMS clarified that once a non-oral version of a previously oralonly drug, such as phosphate binders and calcimimetics, is approved by the Food and Drug Administration ("FDA"), such drug will cease to be considered oral-only. At such time, CMS will commence a process to issue billing codes so that both the oral and nonoral versions of the drug are billable under Part B for a period of at least two years using a transition drug add-on payment adjustment such as average sales price plus 6%, or some other mechanism set in accordance with Section 1847A of the Social Security Act. During this transition period, CMS will not pay outlier payments for these drugs, but will collect data reflecting utilization of both the oral and injectable or intravenous forms of the drugs, as well as payment patterns, in order to more accurately determine the appropriate payment rate to be included in the ESRD PPS for these drugs. At the end of this transition period, CMS will add payment for the oral and non-oral versions of the drug into the ESRD PPS through public rulemaking process similar to that used to set annual ESRD PPS rates. Any failure by CMS to provide adequate additional reimbursement for new drugs or other products that are added to the ESRD PPS could have a material adverse effect on our health care services business and results of operations.

Recent CMS ESRD PPS Payment Rates

On November 6, 2015, CMS published the final ruling regarding the ESRD PPS rate for 2016. We and other large dialysis organizations will experience a 0.2% increase in payments. The base rate per treatment is \$230.39, which represents an approximate reduction of 4%, net, from the 2015 base rate. The 2016 final ruling reflects a net market basket increase of 0.15% (2% less 1.25% PAMA reduction and 0.6% productivity adjustment), application of a wage index budget-neutrality adjustment factor of 1.000495 and application of a refinement budget-neutrality adjustment factor of 0.960319. However, the approximate 4% reduction is almost completely offset with CMS proposed case mix adjustments based upon their analysis of the fiscal years 2012 and 2013.

On June 30, 2016, CMS published a proposed ruling regarding the ESRD PPS rate for 2017. We and other large dialysis organizations will experience a 0.3% increase in payments as compared to the ESRD PPS rate for 2016. The proposed base rate per treatment is \$231.04, which reflects a reduced market basket increase of 0.35%,

application of a wage index budget neutrality adjustment factor of 1.0004482 and application of budget-neutrality adjustment factors of 0.999552 and 0.999729. This proposed rule also includes potential changes to the 2019 and 2020 ESRD QIP measures.

Reimbursement Expectation

As a consequence of the pressure to decrease health care costs, government reimbursement rate increases in the U.S. have historically been limited and are expected to remain stable in the future. We have generally experienced stable reimbursement globally, including the balancing of unfavorable reimbursement changes in certain countries with favorable changes in other countries. In the future we expect to experience generally stable reimbursements for dialysis services globally. However, any significant decreases in Medicare reimbursement rates could have material adverse effects on our health care services business and, because the demand for dialysis products is affected by Medicare reimbursement, on our products business. To the extent that increases in operating costs that are affected by inflation, such as labor and supply costs, are not fully reflected in a compensating increase in reimbursement rates, our business and results of operations may be adversely affected.

Participation in new Medicare Payment Arrangements

CMS is working with various health care providers to develop, refine and implement innovative models of care for Medicare and Medicaid beneficiaries. The extent to which the long-term operation and evolution of these care models, including Accountable Care Organizations, Bundled Payments for Care Improvement Initiative, the Comprehensive ESRD Care Model (which includes the development of ESRD Seamless Care Organizations), and other models, will impact the health care market over time is uncertain. Our U.S. health care provider businesses may choose to participate in certain of these models in certain markets either as a partner with other providers or independently. As existing and new models of care emerge and evolve both in the government and private sectors, patients may choose to be treated by or may be assigned by CMS or other insurers to another provider's care organization, which could have a materially adverse effect on our revenues, earnings and cash flow. As discussed below, we are currently participating in certain of these models.

We participate in CMS's Comprehensive ESRD Care Model ("the CEC Model"), through ESRD Seamless Care Organizations ("ESCOs") in six markets. The CEC Model seeks to deliver better health outcomes for ESRD patients while lowering Medicare's costs. ESCOs that achieve the program's minimum quality thresholds and generate reductions in CMS's cost of care above certain thresholds for the ESRD patients covered by the ESCO will receive a share of the cost savings. Our ESCOs also share in the risk of cost increases above certain thresholds and are required to reimburse CMS a share of any such increases. The CEC Model commenced on October 1, 2015, and the initial agreement period lasts three years. Thereafter, CMS may offer to extend an ESCO's agreement for an additional two years based on the ESCO's performance.

The Bundled Payments for Care Improvement ("BPCI") initiative is a CMS three-year pilot initiative involving bundled payments for the individual services, including acute inpatient hospital services, physician services, and post-acute services, furnished to Medicare beneficiaries during a single episode of illness or course of treatment. Our majority-owned subsidiary, Sound Inpatient Physicians, Inc. ("Sound") commenced participation under BPCI in April 2015 in several markets. Under the BPCI, Sound has the ability to receive additional payments if its physicians are able to deliver quality care at a cost that is lower than certain established benchmarks, but it also has the risk of incurring financial penalties if it is unsuccessful. Should Sound fail to perform as required under its BPCI agreement, CMS may terminate Sound's participation in the BPCI program, in whole or in part.

We have entered into various arrangements with both government and private sector health care insurers which involve taking risk for the complete care of certain ESRD patients in exchange for set payments. We are currently operating Medicare Advantage ESRD Chronic Special Needs Plan ("MA-CSNP") in three states. MA-CSNPs are Medicare Advantage health plans offered by private companies that contract with Medicare to provide patients with Medicare benefits. Enrollment in these plans is limited to special needs individuals with specific severe or disabling chronic conditions, such as ESRD. Our MA-CSNPs provide services, including Care Coordination services, and receive capitated payments from Medicare for the complete care of enrolled ESRD patients. On April 4, 2016, CMS finalized the 2017 payments for Medicare Advantage plans and the Part D Prescription Drug Program. CMS expects a revenue change of 0.85% without consideration for expected growth in coding acuity which typically provides an additional 2.2%.

We also participate in sub-capitation and other shared savings arrangements with certain Medicare Advantage plans and Accountable Care organizations under which we assume risk in providing care to the plans' ESRD patients while paid on a per patient per month basis.

Company Structure

Our operating segments are the North America Segment, the EMEA Segment, the Asia-Pacific Segment and the Latin America Segment. Our management evaluates each segment using measures that reflect all of the segment's controllable revenues and expenses. With respect to the performance of business operations, our management believes that the most appropriate U.S. GAAP measures are revenue, operating income and operating income margin. We do not include income taxes as we believe this is outside the segments' control. Financing is a corporate function which our segments do not control. Therefore, we do not include interest expense relating to financing as a segment measurement. Similarly, we do not allocate certain costs which relate primarily to certain headquarter overhead charges, including accounting and finance, because we believe that these costs are also not within the control of the individual segments. Production of products, production asset management, quality management and procurement related to production are centrally managed at Corporate. The Company's global research and development is also centrally managed at Corporate. These Corporate activities do not fulfill the definition of a segment. Products are transferred to the segments at cost; therefore no internal profit is generated. The associated internal revenues for the product transfers and their elimination are recorded as Corporate activities (See Note 13 in this report). Capital expenditures for production are based on the expected demand of the segments and consolidated profitability considerations. In addition, certain revenues, investments and intangible assets, as well as any related expenses, are not allocated to a segment but accounted for as Corporate, Accordingly, all of these items are excluded from our analysis of segment results and are discussed below in our consolidated results of operations.

Results of Operations

The following tables summarize our financial performance and certain operating results by principal reporting segment and Corporate for the periods indicated. 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. See the table below:

	For the three months ended June 30,			For the six months ended June 30,				
		2016		2015		2016		2015
		(in m	illior	ns)		(in mi	llions	5)
Total revenue (1)								
North America	\$	3,168	\$	2,946	\$	6,212	\$	5,717
EMEA		676		668		1,307		1,297
Asia-Pacific		397		376		771		729
Latin America		175		203		328		401
Corporate		4		6	-	8		15
Total		4,420		4,199		8,626		8,159
Operating income								
North America		513		428		949		768
EMEA		139		134		269		275
Asia-Pacific		75		67		140		152
Latin America		16		16		27		34
Corporate		(102)		(98)		(204)		(178)
Total		641		547		1,181		1,051
Interest income		17		13		28		73
Interest expense		(119)		(115)		(236)		(277)
Income tax expense		(169)		(135)		(306)		(273)
Net Income		370		310		667		574
Less: Net Income attributable to noncontrolling interests		(76)		(69)		(145)		(124)
Net Income attributable to	-	(70)		(09)		(143)		(124)
shareholders of FMC-AG & Co.								
KGaA	\$	294	\$	241	\$	522	\$	450

⁽¹⁾ Net of patient service bad debt provision

Three months ended June 30, 2016 compared to three months ended June 30, 2015

Consolidated Financials

Key Indicators for Consolidated Financial Statements

	For the three months ended		Cha	ange in %
	June 30,			at Constant
	2016	2015	as reported	Exchange Rates(1)
Revenue in \$ million ⁽²)	4,420	4,199	5%	7%
Health Care ⁽²⁾	3,571	3,345	7%	8%
Dialysis Products	849	854	(1%)	2%
Number of dialysis treatments Same market treatment growth in	11,547,779	11,136,497	4%	
%	3.1%	4.3%		
Gross profit as a % of revenue	31.6%	30.9%		
Selling, general and administrative costs as a % of revenue	16.6%	17.2%		
Operating income in \$ million	641	547	17%	
Operating income margin in %	14.5%	13.0%		
Delivered EBIT in \$ million ⁽³⁾ Net income attributable to shareholders of FMC-AG & Co.	565	478	18%	
KGaA in \$ million	294	241	22%	
Basic earnings per share in \$	0.96	0.79	22%	

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation - Constant Currency" below.

Total Revenue increased by 5% (7% increase at Constant Exchange Rates) to \$4,420 million for the three months ended June 30, 2016 from \$4,199 million in the same period of 2015 due to an increase in Health Care revenue.

Health Care revenue increased by 7% to \$3,571 million (8% increase at Constant Exchange Rates) for the three months ended June 30, 2016 from \$3,345 million in the same period of 2015, mainly due to increases in organic revenue per treatment (4%), growth in same market treatments (3%), and contributions from acquisitions (2%), partially offset by the negative effect of exchange rate fluctuations (1%) and by the effect of closed or sold clinics (1%).

Dialysis treatments increased by 4% for the three months ended June 30, 2016 as compared to the same period in 2015. The increase is due to same market treatment growth (3%), and contributions from acquisitions (2%) partially offset by the effect of closed or sold clinics (1%).

At June 30, 2016, we owned, operated or managed (excluding those managed but not consolidated in the U.S.) 3,504 dialysis clinics compared to 3,421 dialysis clinics at June 30, 2015. During the three months ended June 30, 2016, we acquired 58 dialysis clinics, opened 27 dialysis clinics and combined or closed 13 clinics. The number of patients

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

treated in dialysis clinics that we own, operate or manage (excluding patients of dialysis clinics managed but not consolidated in the U.S.) increased by 4% to 301,548 at June 30, 2016 from 290,658 at June 30, 2015.

Dialysis product revenue decreased by 1% (2% increase at Constant Exchange Rates) to \$849 million as compared to \$854 million in the same period of 2015. The increase at Constant Exchange Rates was driven by increased sales of dialyzers, machines, peritoneal dialysis products, bloodlines as well as hemodialysis solutions and concentrates, partially offset by lower sales of renal pharmaceuticals.

The increase in gross profit margin to 31.6% from 30.9% primarily reflects increases in the North America Segment and the Asia-Pacific Segment. The increase in the North America Segment was mainly due to lower costs for health care supplies and a favorable impact from higher volume with commercial payors, partially offset by higher personnel expense related to dialysis services and an unfavorable impact from pharmacy services as a result of price increases for oral medications. The increase in the Asia-Pacific Segment was driven by business growth and favorable foreign exchange effects.

Selling, general and administrative ("SG&A") expenses increased to \$732 million in the three months ended June 30, 2016 from \$723 million in the same period of 2015. SG&A expenses as a percentage of sales decreased to 16.6% for the three months ended June 30, 2016 in comparison with 17.2% in the same period of 2015 due to decreases in the North America Segment and the EMEA Segment. The decrease in the North America Segment was attributable to a favorable impact from endovascular and cardiovascular services related to a gain from a divestiture as well as a gain from the collection of a purchase price escrow claim, lower legal expenses and the impact from increased sales volumes related to pharmacy services, partially offset by an unfavorable impact from hospitalist and intensivist services due to infrastructure development. The decrease in the EMEA Segment was driven by a favorable foreign exchange effects and the impact from increased sales.

Research and development ("R&D") expenses increased by 12% to \$39 million for the three months ended June 30, 2016 from \$34 million for the same period of 2015 driven by higher personnel expense.

Income from equity method investees increased to \$13 million for the three months ended June 30, 2016 from \$7 million for the same period of 2015. This increase is primarily related to higher income from the Vifor Fresenius Medical Care Renal Pharma Ltd. joint venture due to increased revenue resulting from the expansion of its product portfolio.

Operating income increased to \$641 million for the three months ended June 30, 2016 from \$547 million for the same period in 2015. Operating income margin increased to 14.5% for the three months ended June 30, 2016 as compared to 13.0% for the same period in 2015 as a result of increased gross profit margin, decreased SG&A as a percentage of revenue and increased income from equity method investees.

Delivered EBIT increased to \$565 million for the three months ended June 30, 2016 from \$478 million for the same period in 2015 as a result of increased operating income, partially offset by increased noncontrolling interests driven by higher operating income of our joint ventures involving dialysis clinics and the creation of new dialysis clinic joint ventures in the North America Segment.

Interest expense increased by 4% to \$119 million for the three months ended June 30, 2016 from \$115 million for the same period in 2015 due to the higher impact of the valuation of the embedded derivative related to the equity- neutral convertible bonds issued in September 2014 and the related call option on our shares. Interest income increased by 32% to \$17 million for the three months ended June 30, 2016 as compared

to \$13 million for the same period in 2015 due to the higher impact of the valuation of the derivative embedded in the convertible debt and the related call option on our shares as well as interest income related to delayed payments, partially offset by lower interest income due to the repayment of interest bearing notes receivables in the fourth quarter of 2015.

Income tax expense increased to \$169 million for the three months ended June 30, 2016 as compared to \$135 million for the same period in 2015. The effective tax rate increased to 31.3% from 30.4% for the same period of 2015 mainly driven by a proportionately lower increase of tax-free income attributable to noncontrolling interests compared to a higher increase in income before taxes.

Net income attributable to noncontrolling interests for the three months ended June 30, 2016 increased to \$76 million from \$69 million for the same period of 2015 primarily driven by higher operating income of joint ventures with dialysis clinics, but at lower margins and, to a lesser extent, the creation of new joint ventures in the North America Segment, partially offset by decreased noncontrolling interest expense related to Care Coordination.

Net income attributable to shareholders of FMC-AG & Co. KGaA for the three months ended June 30, 2016 increased by 22% to \$294 million from \$241 million for the same period in 2015 as a result of the combined effects of the items discussed above.

Basic earnings per share increased by 22% for the three months ended June 30, 2016 to \$0.96 as compared with \$0.79 for the same period in 2015 primarily due to the increase in net income attributable to shareholders of FMC-AG & Co. KGaA described above. The average weighted number of shares outstanding for the period was approximately 305.5 million in 2016 (304.2 million in 2015).

We employed 106,556 people (full-time equivalents) as of June 30, 2016 compared to 102,893 as of June 30, 2015, an increase of 4%, primarily due to overall growth in our business.

The following discussions pertain to the North America Segment, the EMEA Segment, the Asia-Pacific Segment and the Latin America Segment and the measures we use to manage these segments.

North America Segment

Key Indicators and Business Metrics for North America Segment

	For the three months ended June 30,		Change in
	2016	2015	%
Total North America Segment			
Revenue in \$ million ⁽¹⁾	3,168	2,946	8%
Health Care ⁽¹⁾	2,938	2,722	8%
Dialysis Products	230	224	2%
Operating income in \$ million	513	428	20%
Operating income margin in %	16.2%	14.5%	
Delivered EBIT in \$ million ⁽²⁾	439	362	21%
Dialysis			
Revenue in \$ million ⁽¹⁾	2,604	2,478	5%
Number of dialysis treatments	7,168,288	6,892,346	4%
Same market treatment growth in %	3.0%	4.2%	
Operating income in \$ million	488	391	25%
Operating income margin in %	18.7%	15.8%	
Delivered EBIT in \$ million ⁽²⁾	422	338	25%
Care Coordination			
Revenue in \$ million ⁽¹⁾	564	468	21%
Operating income in \$ million	25	37	(32%)
Operating income margin in %	4.4%	7.8%	
Delivered EBIT in \$ million ⁽²⁾ Member Months Under Medical Cost	17	24	(31%)
Management ^{(3),(4)}	91,392	40,287	127%
Medical Cost Under Management in \$ million ^{(3),(4)}	658	432	52%
Care Coordination Patient Encounters (3),(4)	1,338,695	1,270,257	5%

⁽¹⁾ Net of patient service bad debt provision.

Dialysis

Revenue

Dialysis revenue increased for the three months ended June 30, 2016 by 5% to \$2,604 million from \$2,478 million in the same period of 2015.

Dialysis care revenue increased for the three months ended June 30, 2016 by 5% to \$2,374 million from \$2,254 million in the same period of 2015. This increase was driven by same market treatment growth (3%), increases in organic revenue per treatment (2%) and contributions from acquisitions (1%), partially offset by the effect of closed or sold clinics (1%).

⁽²⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below

⁽³⁾ For further information on these metrics, please refer to the discussion below of our Care Coordination measures under "Care Coordination Business Metrics for Presentation."

⁽⁴⁾ The 2016 metric may be understated due to a physician mapping issue related to the BPCI program within a CMS system which has not yet been resolved. Additionally, data presented for the metrics are subject to finalization by CMS, which may result in changes from previously reported metrics.

Dialysis treatments increased by 4% for the three months ended June 30, 2016 as compared to the same period in 2015 primarily due to same market treatment growth (3%) and contributions from acquisitions (1%). At June 30, 2016, 186,096 patients (a 4% increase from June 30, 2015) were being treated in the 2,249 dialysis clinics that we own or operate in the North America Segment, compared to 178,766 patients treated in 2,205 dialysis clinics at June 30, 2015.

In the U.S., the average revenue per treatment was \$352 for the three months ended June 30, 2016 and \$346 for the same period in 2015. The increase was mainly attributable to a favorable impact from higher volume with commercial payors.

Cost per treatment in the U.S. decreased to \$282 for the three months ended June 30, 2016 from \$286 in the same period of 2015. This decrease was largely driven by a favorable impact from lower cost for health care supplies, partially offset by higher personnel expense.

Dialysis product revenue increased by 2% to \$230 million for the three months ended June 30, 2016 as compared to \$224 million in the same period in 2015. This was driven by higher sales of machines, dialyzers and peritoneal dialysis products, partially offset by lower sales of renal pharmaceuticals.

Operating Income

Dialysis operating income increased to \$488 million for the three months ended June 30, 2016 as compared to \$391 million in the same period in 2015. Operating income margin increased to 18.7% for the three months ended June 30, 2016 from 15.8% for the same period in 2015, due to lower costs from health care supplies, a favorable impact from higher volume with commercial payors, lower legal expenses and increased income from equity method investees, partially offset by higher personnel expense.

Delivered EBIT

Dialysis delivered EBIT increased by 25% to \$422 million for the three months ended June 30, 2016 from \$338 million for the same period of 2015 mainly as the result of increased operating income, partially offset by increased noncontrolling interests driven by higher operating income of our joint ventures involving dialysis clinics, but at lower margins and, to a lesser extent, the creation of new joint ventures.

Care Coordination

Revenue

Care Coordination revenue increased by 21% to \$564 million for the three months ended June 30, 2016 from \$468 million for the same period of 2015. This increase was driven by increases in organic revenue growth (17%), contributions from acquisitions (3%) and reduction of bad debt (1%).

Operating Income

Care Coordination operating income decreased to \$25 million for the three months ended June 30, 2016 from \$37 million for the same period of 2015. The operating income margin decreased to 4.4% for the three months ended June 30, 2016 from 7.8% mainly driven by increased costs for hospitalist and intensivist services due to infrastructure development, growth in lower margin health plan services as well as higher costs for supplies for laboratory services, partially offset by a favorable impact from endovascular and cardiovascular services related to a gain from a divestiture as well as a gain from the collection of a purchase price escrow claim.

Delivered EBIT

Care Coordination delivered EBIT decreased to \$17 million for the three months ended June 30, 2016 from \$24 million for the same period of 2015 mainly as the result of decreased operating income partially offset by decreased noncontrolling interests effects.

Member Months Under Medical Cost Management

Care Coordination's member months under medical cost management for the three months ended June 30, 2016 was 91,392 months as compared to 40,287 months for the same period of 2015. The increase in membership volume was due to BPCI development, the commencement of ESCOs and inclusion of ESCO amounts in the fourth quarter of 2015 as well as the contribution from MA-CSNPs in the first quarter of 2016. See note 4 to the table "Key Indicators and Business Metrics for North America Segment," above.

Medical Cost Under Management

Care Coordination's medical cost under management for the three months ended June 30, 2016 was \$658 million as compared to \$432 million for the same period of 2015. The increase in medical cost under management was attributable to the commencement of ESCOs and inclusion of ESCO amounts in the fourth quarter of 2015, BPCI development as well as the contribution from MA-CSNPs in the first quarter of 2016. See note 4 to the table "Key Indicators and Business Metrics for North America Segment," above.

Care Coordination Patient Encounters

Care Coordination's patient encounters for the three months ended June 30, 2016 were 1,338,695 encounters and procedures as compared to 1,270,257 encounters and procedures for the three months ended June 30, 2015. The increase was driven by patient encounters and procedures provided by Fresenius Medical Care Rx Bone Mineral Metabolism ("Rx BMM") program, hospitalist and intensivist services, urgent care centers, vascular procedures as well as cardiovascular and endovascular services. See note 4 to the table "Key Indicators and Business Metrics for North America Segment," above.

EMEA Segment

Key Indicators for EMEA Segment

			Change in %		
	For the thro end June	led		at Constant Exchange	
	2016	2015	as reported	Rates ⁽¹⁾	
Revenue in \$ million ⁽²⁾	676	668	1%	3%	
Health Care ⁽²⁾	331	309	7%	9%	
Dialysis Products	345	359	(4%)	(3%)	
Number of dialysis treatments	2,217,107	2,034,186	9%		
Same market treatment growth in %	3.4%	3.8%			
Operating income in \$ million	139	134	4%		
Operating income margin in %	20.6%	20.1%			
Delivered EBIT in \$ million ⁽³⁾	139	133	4%		

- (1) For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation-Constant Currency" below.
- (2) Net of patient service bad debt provision.
- (3) For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation Delivered EBIT" below.

Revenue

Total revenue for the EMEA Segment increased by 1% (3% increase at Constant Exchange Rates) to \$676 million for the three months ended June 30, 2016 as compared to \$668 million for the same period of 2015. Health care service revenue for the EMEA Segment increased by 7% (9% increase at Constant Exchange Rates) to \$331 million during the three months ended June 30, 2016 as compared to \$309 million for the same period of 2015. This is a result of contributions from acquisitions (7%) and increases in organic revenue growth per treatment (3%), partially offset by the negative effect of exchange rate fluctuations (2%) and the effect of closed or sold clinics (1%). Dialysis treatments increased by 9% for the three months ended June 30, 2016 over the same period in 2015 mainly due to contributions from acquisitions (7%) and same market treatment growth (3%), partially offset by the effect of closed or sold clinics (1%). As of June 30, 2016, we had 58,528 patients (9% increase from June 30, 2015) being treated at the 700 dialysis clinics that we own, operate or manage in the EMEA Segment compared to 53,546 patients treated at 648 clinics at June 30, 2015.

Dialysis product revenue for the three months ended June 30, 2016 decreased by 4% (3% decrease at Constant Exchange Rates) to \$345 million as compared to \$359 million in the same period of 2015. The decrease at Constant Exchange Rates was driven by lower sales of dialyzers, machines, renal pharmaceuticals and bloodlines, partially offset by higher sales of products for acute care treatments and peritoneal dialysis products.

Operating Income

Operating income increased to \$139 million for the three months ended June 30, 2016 as compared to \$134 million for the same period in 2015. Operating income margin increased to 20.6% for the three months ended June 30, 2016 from 20.1% for the same period in 2015 mainly due to favorable foreign exchange effects and a favorable impact from manufacturing driven by higher volumes and production efficiencies, partially offset by lower income from equity method investees.

Delivered EBIT

Delivered EBIT increased by 4% to \$139 million for the three months ended June 30, 2016 as compared to \$133 million for the same period in 2015 due to increased operating income, partially offset by increased noncontrolling interests.

Asia-Pacific Segment

Key Indicators for Asia-Pacific Segment

			Change in %		
	For the three months ended June 30,		as reported	at Constant Exchange Rates ⁽¹⁾	
Revenue in \$ million ⁽²⁾	397	376	5%	6%	
•					
Health Care ⁽²⁾	177	164	8%	2%	
Dialysis Products	220	212	4%	9%	
Number of dialysis treatments Same market treatment growth in	978,819	942,855	4%		
%	5.0%	3.3%			
Operating income in \$ million	75	67	12%		
Operating income margin in %	18.9%	17.8%			
Delivered EBIT in \$ million ⁽³⁾	73	65	13%		

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation-Constant Currency" below.

Revenue

Total revenue for the Asia-Pacific Segment increased by 5% (6% increase at Constant Exchange Rates) to \$397 million for the three months ended June 30, 2016 as compared to \$376 million for the same period of 2015. Health care service revenue for the Asia-Pacific Segment increased during the three months ended June 30, 2016 by 8% (2% increase at Constant Exchange Rates) to \$177 million from \$164 million in the same period of 2015. This increase is a result of the effect of exchange rate fluctuations (6%) same market treatment growth (5%), partially offset by the effect of closed or sold clinics (2%) and decreases in organic revenue growth per treatment (1%). Dialysis treatments increased by 4% for the three months ended June 30, 2016 over the same period in 2015 mainly due to same market treatment growth (5%), partially offset by the effect of closed or sold clinics (1%). As of June 30, 2016, we had 27,007 patients (a 4% increase from June 30, 2015) being treated at the 324 dialysis clinics that we own, operate or manage in the Asia-Pacific Segment compared to 26,024 patients treated at 320 clinics at June 30, 2015.

Dialysis product revenue for the three months ended June 30, 2016 increased by4% (9% increase at Constant Exchange Rates) to \$220 million compared to \$212 million in the same period of 2015. The increase at Constant Exchange Rates was driven by increased sales of dialyzers, bloodlines, peritoneal dialysis products, machines and hemodialysis solutions and concentrates.

Operating Income

Operating income increased by 12% to \$75 million for the three months ended June 30, 2016 as compared to \$67 million for the same period in 2015. Operating income

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see " Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

margin increased to 18.9% for the three months ended June 30, 2016 compared to 17.8% in the same period of 2015 due to favorable foreign exchange effects and business growth, partially offset by lower income from equity method investees.

Delivered EBIT

Delivered EBIT increased by 13% to \$73 million for the three months ended June 30, 2016 as compared to \$65 million for the same period in 2015 due to increased operating income with virtually no change in noncontrolling interests.

Latin America Segment

Key Indicators for Latin America Segment

•			Change in %		
	For the three months ended June 30,		as	at Constant Exchange	
	2016	2015	reported	Rates ⁽¹⁾	
Revenue in \$ million ⁽²⁾	175	203	(14%)	9%	
Health Care ⁽²⁾	125	150	(17%)	9%	
Dialysis Products	50	53	(5%)	8%	
Number of dialysis treatments	1,183,565	1,267,110	(7%)		
Same market treatment growth in %	1.5%	6.8%			
Operating income in \$ million	16	16	4%		
Operating income margin in %	9.3%	7.8%			
Delivered EBIT in \$ million ⁽³⁾	16	16	5%		

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation-Constant Currency" below.

Revenue

Total revenue for the Latin America Segment decreased by 14% (9% increase at Constant Exchange Rates) to \$175 million for the three months ended June 30, 2016 as compared to \$203 million for the same period of 2015. Health care service revenue for the Latin America Segment decreased by 17% (9% increase at Constant Exchange Rates) during the three months ended June 30, 2016 to \$125 million as compared to \$150 million in the same period of 2015. This decrease is a result of the negative effect of exchange rate fluctuations (26%) and the effect of closed or sold clinics (mainly in Venezuela) (12%), partially offset by increases in organic revenue per treatment (17%), growth in same market treatments (2%) and contributions from acquisitions (2%). Dialysis treatments decreased by 7% for the three months ended June 30, 2016 over the same period in 2015 mainly due to the effect of closed or sold clinics (mainly in Venezuela) (10%), partially offset by same market treatment growth (2%) and contributions from acquisitions (1%). As of June 30, 2016, we had 29,917 patients (a 7%) decrease from June 30, 2015) being treated at the 231 dialysis clinics that we own, operate or manage in the Latin America Segment compared to 32,322 patients treated at 248 clinics at June 30, 2015.

Dialysis product revenue for the three months ended June 30, 2016 decreased by 5% (8% increase at Constant Exchange Rates) to \$50 million compared to \$53 million in the same period of 2015. The 8% increase at Constant Exchange Rates was mainly driven by

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

higher sales of dialyzers, hemodialysis solutions and concentrates, machines and bloodlines, partially offset by lower sales of peritoneal dialysis products.

Operating Income

Operating income remained stable at \$16 million for the three months ended June 30, 2016 as compared to the same period in 2015 (this represents a 4% increase without effect from rounding). Operating income margin increased to 9.3% for the three months ended June 30, 2016 from 7.8% for the same period in 2015 mainly due to favorable foreign exchange effects and the impact from higher revenue in the region at Constant Exchange Rates, partially offset by higher costs related to inflation.

Delivered EBIT

Delivered EBIT remained stable at \$16 million for the three months ended June 30, 2016 as compared to the same period in 2015 (this represents a 5% increase without effect from rounding due to increased operating income with virtually no change in noncontrolling interests).

Six months ended June 30, 2016 compared to six months ended June 30, 2015

Consolidated Financials

Key Indicators for Consolidated Financial Statements

	For the six months ended		Cha	ange in %
<u>-</u>	June 30,		as	at Constant Exchange
<u>-</u>	2016	2015	reported	Rates(1)
Revenue in \$ million ⁽²)	8,626	8,159	6%	8%
Health Care(2)	6,985	6,527	7%	9%
Dialysis Products	1,641	1,632	1%	4%
Number of dialysis treatments	22,821,121	21,907,899	4%	
Same market treatment growth in %	3.5%	4.2%		
Gross profit as a % of revenue	31.5%	30.4%		
Selling, general and administrative costs as a % of revenue	17.3%	16.9%		
Operating income in \$ million	1,181	1,051	12%	
Operating income margin in %	13.7%	12.9%		
Delivered EBIT in \$ million ⁽³⁾ Net income attributable to shareholders of FMC-AG & Co.	1,036	927	12%	
KGaA in \$ million	522	450	16%	
Basic earnings per share in \$	1.71	1.48	15%	

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation - Constant Currency" below.

Total Revenue increased by 6% (8% increase at Constant Exchange Rates) to \$8,626 million for the six months ended June 30, 2016 from \$8,159 million in the same period of 2015 due to increases in Health Care revenue and dialysis product revenue.

Health Care revenue increased by 7% to \$6,985 million (9% increase at Constant Exchange Rates) for the six months ended June 30, 2016 from \$6,527 million in the same period of 2015, mainly due to growth in same market treatments (4%), increases in organic revenue per treatment (3%), an increase in dialysis days (1%) and contributions from acquisitions (1%), partially offset by the negative effect of exchange rate fluctuations (2%).

Dialysis treatments increased by 4% for the six months ended June 30, 2016 as compared to the same period in 2015. The increase is due to same market treatment growth (4%), an increase in dialysis days (1%) and contributions from acquisitions (1%) partially offset by the effect of closed or sold clinics (2%).

Dialysis product revenue increased by 1% (4% increase at Constant Exchange Rates) to \$1,641 million as compared to \$1,632 million in the same period of 2015. The increase at Constant Exchange Rates was driven by increased sales of dialyzers, machines, bloodlines, peritoneal dialysis products, hemodialysis solutions and concentrates and products for acute care treatments, partially offset by lower sales of renal pharmaceuticals.

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

The increase in gross profit margin to 31.5% from 30.4% primarily reflects increases in the North America Segment, the Asia-Pacific Segment and the EMEA Segment. The increase in the North America Segment was mainly due to lower costs for health care supplies and a favorable impact from higher volume with commercial payors, partially offset by higher personnel expense related to dialysis services, an unfavorable impact from pharmacy services as a result of price increases in oral medications and growth in health plan services at lower than average margins. The increase in the Asia-Pacific Segment was driven by favorable foreign exchange effects. The increase in the EMEA Segment was primarily driven by a favorable impact from manufacturing related to higher volumes and production efficiencies, partially offset by unfavorable foreign exchange effects.

Selling, general and administrative ("SG&A") expenses increased to \$1,491 million in the six months ended June 30, 2016 from \$ 1,378 million in the same period of 2015. SG&A expenses as a percentage of sales increased to 17.3% for the six months ended June 30, 2016 in comparison with 16.9% in the same period of 2015 due to increases in the Asia-Pacific Segment, Corporate and the EMEA Segment, partially offset by a decrease in the North America Segment. The increase in the Asia-Pacific Segment was mainly due to unfavorable foreign exchange effects, costs associated with changes in the Management Board and increased costs related to furthered sales development. The increase at Corporate was primarily driven by higher legal and consulting expenses related to compliance investigations we are conducting (see Note 10in this report) and higher overhead costs for manufacturing. The increase in the EMEA Segment was driven by unfavorable foreign exchange effects and higher bad debt expense, partially offset by the impact from higher sales. The decrease in the North America Segment was mainly due to lower legal expense, the impact from higher sales related to pharmacy services and a favorable impact from cardiovascular and endovascular services related to a gain from a divestiture as well as a gain from the collection of a purchase price escrow claim, partially offset by an unfavorable impact from hospitalist and intensivist services due to infrastructure development.

Research and development ("R&D") expenses increased by 17% to \$76 million for the six months ended June 30, 2016 from \$65 million for the same period of 2015 driven by higher personnel expense.

Income from equity method investees increased to \$32 million for the six months ended June 30, 2016 from \$13 million for the same period of 2015. This increase is primarily related to higher income from the Vifor Fresenius Medical Care Renal Pharma Ltd. joint venture due to increased revenue resulting from the expansion of its product portfolio.

Operating income increased to \$1,181 million for the six months ended June 30, 2016 from \$1,051 million for the same period in 2015. Operating income margin increased to 13.7% for the six months ended June 30, 2016 as compared to 12.9% for the same period in 2015 as a result of increased gross profit margin and income from equity method investees, partially offset by an increase in SG&A as a percentage of revenue.

Delivered EBIT increased to \$1,036 million for the six months ended June 30, 2016 from \$927 million for the same period in 2015 as a result of increased operating income.

Interest expense decreased by 15% to \$236 million for the six months ended June 30, 2016 from \$277 million for the same period in 2015 due to the lower impact of the valuation of the embedded derivative related to the equity- neutral convertible bonds issued in September 2014 and the related call option on our shares. Interest income decreased by 61% to \$28 million for the six months ended June 30, 2016 as compared to \$73 million for the same period in 2015 due to the lower impact of the valuation of the

derivative embedded in the convertible debt and the related call option on our shares as well as the repayment of interest bearing notes receivables in the fourth quarter of 2015.

Income tax expense increased to \$306 million for the six months ended June 30, 2016 as compared to \$273 million for the same period in 2015. The effective tax rate decreased to 31.5% from 32.2% for the same period of 2015 mainly driven by higher tax-free income from equity method investees and increased tax-free income attributable to noncontrolling interests.

Net income attributable to noncontrolling interests for the six months ended June 30, 2016 increased to \$145 million from \$124 million for the same period of 2015 primarily driven by higher operating income of joint ventures with dialysis clinics, but at lower margins and, to a lesser extent, the creation of new dialysis clinic joint ventures in the North America Segment, partially offset by decreased noncontrolling interest expense related to Care Coordination.

Net income attributable to shareholders of FMC-AG & Co. KGaA for the six months ended June 30, 2016 increased by 16% to \$522 million from \$450 million for the same period in 2015 as a result of the combined effects of the items discussed above.

Basic earnings per share increased by 15% for the six months ended June 30, 2016 to \$1.71 as compared with \$1.48 for the same period in 2015 primarily due to the increase in net income attributable to shareholders of FMC-AG & Co. KGaA described above. The average weighted number of shares outstanding for the period was approximately 305.4 million in 2016 (303.9 million in 2015).

The following discussions pertain to the North America Segment, the EMEA Segment, the Asia-Pacific Segment and the Latin America Segment and the measures we use to manage these segments.

North America Segment

Key Indicators and Business Metrics for North America Segment

	For the six months ended June 30,		Change in
	2016	2015	%
Total North America Segment			
Revenue in \$ million ⁽¹⁾	6,212	5,717	9%
Health Care ⁽¹⁾	5,770	5,293	9%
Dialysis Products	442	424	4%
Operating income in \$ million	949	768	24%
Operating income margin in %	15.3%	13.4%	
Delivered EBIT in \$ million ⁽²⁾	809	649	25%
Dialysis			
Revenue in \$ million ⁽¹⁾	5,126	4,815	6%
Number of dialysis treatments	14,221,402	13,527,268	5%
Same market treatment growth in %	3.5%	4.0%	
Operating income in \$ million	914	716	28%
Operating income margin in %	17.8%	14.9%	
Delivered EBIT in \$ million ⁽²⁾	791	619	28%
Care Coordination			
Revenue in \$ million ⁽¹⁾	1,086	902	20%
Operating income in \$ million	35	52	(32%)
Operating income margin in %	3.2%	5.8%	
Delivered EBIT in \$ million ⁽²⁾ Member Months Under Medical Cost	18	30	(39%)
Management ^{(3),(4)}	184,767	44,592	314%
Medical Cost Under Management in \$ million ^{(3),(4)}	1,318	463	185%
Care Coordination Patient Encounters (3),(4)	2,645,771	2,542,304	4%

⁽¹⁾ Net of patient service bad debt provision.

Dialysis

Revenue

Dialysis revenue increased for the six months ended June 30, 2016 by 6% to \$5,126 million from \$4,815 million in the same period of 2015.

Dialysis care revenue increased for the six months ended June 30, 2016 by 7% to \$4,684 million from \$4,391 million in the same period of 2015. This increase was driven by same market treatment growth (3%), increases in organic revenue per treatment (2%), an increase in dialysis days (1%) and contributions from acquisitions (1%).

⁽²⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below

⁽³⁾ For further information on these metrics, please refer to the discussion below of our Care Coordination measures under "Care Coordination Business Metrics for Presentation."

⁽⁴⁾ The 2016 metric may be understated due to a physician mapping issue related to the BPCI program within a CMS system which has not yet been resolved. Additionally, data presented for the metrics are subject to finalization by CMS, which may result in changes from previously reported metrics.

Dialysis treatments increased by 5% for the six months ended June 30, 2016 as compared to the same period in 2015 primarily due to same market treatment growth (3%), an increase in dialysis days (1%) and contributions from acquisitions (1%).

In the U.S., the average revenue per treatment was \$350 for the six months ended June 30, 2016 and \$344 for the same period in 2015. The increase was mainly attributable to a favorable impact from higher volume with commercial payors.

Cost per treatment in the U.S. decreased to \$282 for the six months ended June 30, 2016 from \$287 in the same period of 2015. This decrease was largely driven by a favorable impact from lower cost for health care supplies, partially offset by higher personnel expense.

Dialysis product revenue increased by 4% to \$442 million for the six months ended June 30, 2016 as compared to \$424 million in the same period in 2015. This was driven by higher sales of machines, dialyzers and peritoneal dialysis products, partially offset by lower sales of renal pharmaceuticals.

Operating Income

Dialysis operating income increased to \$914 million for the six months ended June 30, 2016 as compared to \$716 million in the same period in 2015. Operating income margin increased to 17.8% for the six months ended June 30, 2016 from 14.9% for the same period in 2015, due to lower costs from health care supplies, a favorable impact from higher volume with commercial payors, higher income from equity method investees and lower legal expenses, partially offset by higher personnel expense.

Delivered EBIT

Dialysis delivered EBIT increased by 28% to \$791 million for the six months ended June 30, 2016 from \$619 million for the same period of 2015 mainly as the result of increased operating income, partially offset by increased noncontrolling interests driven by higher operating income of our joint ventures involving dialysis clinics, but at lower margins and, to a lesser extent, the creation of new joint ventures.

Care Coordination

Revenue

Care Coordination revenue increased by 20% to \$1,086 million for the six months ended June 30, 2016 from \$902 million for the same period of 2015. This increase was driven by increases in organic revenue growth (17%), contributions from acquisitions (2%) and reduction of bad debt (1%).

Operating Income

Care Coordination operating income decreased to \$35 million for the six months ended June 30, 2016 from \$52 million for the same period of 2015. The operating income margin decreased to 3.2% for the six months ended June 30, 2016 from 5.8% mainly driven by increased costs for hospitalist and intensivist services due to infrastructure development as well as growth in lower margin health plan and urgent care services and higher costs for supplies for laboratory services, partially offset by a favorable impact from endovascular and cardiovascular services related to a gain from a divestiture as well as a gain from the collection of a purchase price escrow claim and the impact from increased sales of pharmacy services.

Delivered EBIT

Care Coordination delivered EBIT decreased to \$18 million for the six months ended June 30, 2016 from \$30 million for the same period of 2015 mainly as the result of

decreased operating income, partially offset by decreased noncontrolling interests effects.

Member Months Under Medical Cost Management

Care Coordination's member months under medical cost management for the six months ended June 30, 2016 was 184,767 months as compared to 44,592 months for the same period of 2015. The increase in membership volume was attributable to BPCI development, the commencement of ESCOs and inclusion of ESCO amounts in the fourth quarter of 2015 as well as the contribution from MA-CSNPs in the first quarter of 2016. See note 4 to the table "Key Indicators and Business Metrics for North America Segment," above.

Medical Cost Under Management

Care Coordination's medical cost under management for the six months ended June 30, 2016 was \$1,318 million as compared to \$463 million for the same period of 2015. The increase in medical cost under management was attributable to the commencement of ESCOs and inclusion of ESCO amounts in the fourth quarter of 2015, BPCI development as well as the contribution from MA-CSNPs in the first quarter of 2016. See note 4 to the table "Key Indicators and Business Metrics for North America Segment," above.

Care Coordination Patient Encounters

Care Coordination's patient encounters for the six months ended June 30, 2016 were 2,645,771 encounters and procedures as compared to 2,542,304 encounters and procedures for the six months ended June 30, 2015. The increase was driven by patient encounters and procedures provided by Rx BMM program, urgent care centers, hospitalist and intensivist services, vascular procedures as well as cardiovascular and endovascular services. See note 4 to the table "Key Indicators and Business Metrics for North America Segment," above.

EMEA Segment

Key Indicators for EMEA Segment

			Change in %	
	For the six months ended June 30,			at Constant Exchange
	2016	2015	as reported	Rates ⁽¹⁾
Revenue in \$ million ⁽²⁾	1,307	1,297	1%	4%
Health Care ⁽²⁾	632	610	4%	8%
Dialysis Products	675	687	(2%)	1%
Number of dialysis treatments	4,312,717	4,023,243	7%	
Same market treatment growth in %	3.6%	4.0%		
Operating income in \$ million	269	275	(2%)	
Operating income margin in %	20.6%	21.2%		
Delivered EBIT in \$ million ⁽³⁾	267	274	(2%)	

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation-Constant Currency" below.

Revenue

Total revenue for the EMEA Segment increased by 1% (4% increase at Constant Exchange Rates) to \$1,307 million for the six months ended June 30, 2016 as compared to \$1,297 million for the same period of 2015. Health care service revenue for the EMEA Segment increased by 4% (8% increase at Constant Exchange Rates) to \$632 million during the six months ended June 30, 2016 as compared to \$610 million for the same period of 2015. This increase is a result of contributions from acquisitions (5%), same market treatment growth (4%), and an increase in dialysis days (1%), partially offset by the negative impact of exchange rate fluctuations (4%), the effect of closed or sold clinics (1%) and decreases in organic revenue growth per treatment (1%). Dialysis treatments increased by 7% for the six months ended June 30, 2016 over the same period in 2015 mainly due to same market treatment growth (4%) and contributions from acquisitions (4%), partially offset by the effect of closed or sold clinics (1%).

Dialysis product revenue for the six months ended June 30, 2016 decreased by 2% (1% increase at Constant Exchange Rates) to \$675 million as compared to \$687 million in the same period of 2015. The increase at Constant Exchange Rates was driven by increased sales of products for acute care treatments, hemodialysis solutions and concentrates, bloodlines as well as peritoneal dialysis products, partially offset by lower sales of renal pharmaceuticals, machines and dialyzers.

Operating Income

Operating income decreased to \$269 million for the three months ended June 30, 2016 as compared to \$275 million for the same period in 2015. Operating income margin decreased to 20.6% for the six months ended June 30, 2016 from 21.2% for the same period in 2015 mainly due to unfavorable foreign exchange effects, partially offset by a favorable impact from manufacturing driven by higher volumes and production efficiencies and furthered sales development at Constant Exchange Rates.

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

Delivered EBIT

Delivered EBIT decreased by 2% to \$267 million for the six months ended June 30, 2016 as compared to \$274 million for the same period in 2015 due to decreased operating income, partially offset by increased noncontrolling interests effects.

Asia-Pacific Segment

Key Indicators for Asia-Pacific Segment

			Change in %		
		For the six months ended June 30,		at Constant Exchange	
	2016	2015	as reported	Rates ⁽¹⁾	
Revenue in \$ million ⁽²⁾	771	729	6%	8%	
Health Care ⁽²⁾	345	328	5%	3%	
Dialysis Products	426	401	6%	12%	
Number of dialysis treatments Same market treatment growth in	1,949,115	1,862,018	5%		
%	5.9%	3.0%			
Operating income in \$ million	140	152	(8%)		
Operating income margin in %	18.2%	20.8%			
Delivered EBIT in \$ million ⁽³⁾	137	148	(7%)		

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation-Constant Currency" below.

Revenue

Total revenue for the Asia-Pacific Segment increased by 6% (8% increase at Constant Exchange Rates) to \$771 million for the six months ended June 30, 2016 as compared to \$729 million for the same period of 2015. Health care service revenue for the Asia-Pacific Segment increased during the six months ended June 30, 2016 by 5% (3% increase at Constant Exchange Rates) to \$345 million from \$328 million in the same period of 2015. This increase is a result of same market treatment growth (6%) and the effect of exchange rate fluctuations (2%), partially offset by decreases in organic revenue growth per treatment (2%), and the effect of closed or sold clinics (1%). Dialysis treatments increased by 5% for the six months ended June 30, 2016 over the same period in 2015 mainly due to same market treatment growth (6%), partially offset by the effect of closed or sold clinics (1%).

Dialysis product revenue for the six months ended June 30, 2016 increased by 6% (12% increase at Constant Exchange Rates) to \$426 million compared to \$401 million in the same period of 2015. The increase at Constant Exchange Rates was driven by increased sales of dialyzers, bloodlines, machines, peritoneal dialysis products and hemodialysis solutions and concentrates.

Operating Income

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see " Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

Operating income decreased by 8% to \$140 million for the six months ended June 30, 2016 as compared to \$152 million for the same period in 2015. Operating income margin decreased to 18.2% for the six months ended June 30, 2016 compared to 20.8% in the same period of 2015 due to costs associated with changes in the Management Board, unfavorable foreign exchange effects, an adverse impact from manufacturing driven by lower volumes of dialyzers and concentrates and increased costs related to furthered sales development.

Delivered EBIT

Delivered EBIT decreased by 7% to \$137 million for the six months ended June 30, 2016 as compared to \$148 million for the same period in 2015 due to decreased operating income with virtually no change in noncontrolling interests.

Latin America Segment

Key Indicators for Latin America Segment

•			Change in %	
	For the six months ended June 30,		as	at Constant Exchange
	2016	2015	reported	Rates ⁽¹⁾
Revenue in \$ million	328	401	(18%)	7%
Health Care ⁽²⁾	238	296	(20%)	9%
Dialysis Products	90	105	(14%)	2%
Number of dialysis treatments	2,337,887	2,495,370	(6%)	
Same market treatment growth in %	1.8%	6.4%		
Operating income in \$ million	27	34	(19%)	
Operating income margin in %	8.3%	8.4%		
Delivered EBIT in \$ million ⁽³⁾	27	34	(19%)	

⁽¹⁾ For further information on Constant Exchange Rates, see "Non-U.S. GAAP Measures for Presentation-Constant Currency" below.

Revenue

Total revenue for the Latin America Segment decreased by 18% (7% increase at Constant Exchange Rates) to \$328 million for the six months ended June 30, 2016 as compared to \$401 million for the same period of 2015. Health care service revenue for the Latin America Segment decreased by 20% (9% increase at Constant Exchange Rates) during the six months ended June 30, 2016 to \$238 million as compared to \$296 million for the same period of 2015. The health care service revenue decreased as a result of the negative effect of exchange rate fluctuations (29%) and the effect of closed or sold clinics (mainly in Venezuela) (10%), partially offset by increases in organic revenue per treatment (15%), growth in same market treatments (2%), contributions from acquisitions (1%) and an increase in dialysis days (1%). Dialysis treatments decreased by 6% for the six months ended June 30, 2016 over the same period in 2015 mainly due to the effect of closed or

⁽²⁾ Net of patient service bad debt provision.

⁽³⁾ For further information on Delivered EBIT, see "Non-U.S. GAAP Measures for Presentation – Delivered EBIT" below.

sold clinics (10%), partially offset by same market treatment growth (2%), contributions from acquisitions (1%) and an increase in dialysis days (1%).

Dialysis product revenue for the six months ended June 30, 2016 decreased by 14% (2% increase at Constant Exchange Rates) to \$90 million compared to \$105 million in the same period of 2015. The 2% increase at Constant Exchange Rates was driven by increased sales of dialyzers and bloodlines, partially offset by lower sales of peritoneal dialysis products and machines.

Operating Income

Operating income decreased by 19% to \$27 million for the six months ended June 30, 2016 as compared to \$34 million for the same period in 2015. Operating income margin decreased to 8.3% for the six months ended June 30, 2016 from 8.4% for the same period in 2015 mainly due to higher costs related to inflation and an unfavorable impact from manufacturing production costs driven by unfavorable foreign exchange effects and higher costs for quality development, partially offset by the impact from higher revenue in the region as well as the impact from prior year lower margin dialysis service business in Venezuela which was subsequently divested in the third quarter of 2015.

Delivered EBIT

Delivered EBIT decreased by 19% to \$27 million for the six months ended June 30, 2016 as compared to \$34 million for the same period in 2015 due to the impacts noted above in operating income with virtually no change in noncontrolling interests.

Liquidity and Capital Resources

Six months ended June 30, 2016 compared to six months ended June 30, 2015

Our primary sources of liquidity are typically cash provided by operating activities, cash provided by short-term debt from third parties and related parties, as well as proceeds from the issuance of long-term debt and equity securities. We require this capital primarily to finance working capital needs, fund acquisitions and joint ventures, develop free-standing renal dialysis clinics and other health care facilities, purchase equipment for existing or new renal dialysis clinics and production sites, repay debt, pay dividends and repurchase shares (see "Net Cash Provided By (Used In) Investing Activities" and "Net Cash Provided By (Used In) Financing Activities" below).

At June 30, 2016, we had cash and cash equivalents of \$723 million. For information regarding utilization and availability of cash under our principal credit facility (the "Amended 2012 Credit Agreement"), see Note 5 in this report.

Net Cash Provided By (Used In) Operating Activities

In the first six months of 2016 and 2015, we generated net cash provided by operating activities of \$857 million and \$832 million, respectively. Cash provided by operating activities is impacted by the profitability of our business, the development of our working capital, principally inventories, receivables and cash outflows that occur due to a number of specific items as discussed below. The increase in 2016 versus 2015 was mainly a result of increased earnings and reduced inventory levels driven by decreased volume of health care

supplies, particularly due to a decrease in erythropoietin-stimulating agents inventory, partially offset by higher income tax payments due to a tax refund in the comparable period of 2015 and increased earnings in the current period as well as the timing of other working capital items.

The profitability of our business depends significantly on reimbursement rates. Approximately 81% of our revenues are generated by providing health care services, a major portion of which is reimbursed by either public health care organizations or private insurers. Legislative changes could affect reimbursement rates for a significant portion of the services we provide, as well as the scope of coverage. A decrease in reimbursement rates or the scope of coverage could have a material adverse effect on our business, financial condition and results of operations and thus on our capacity to generate cash flow.

Our working capital, which is defined as current assets less current liabilities, was \$2,358 million at June 30, 2016 which decreased from \$2,619 million at December 31, 2015. The change is primarily the result of increased short-term debt due to issuance of short-term notes under our commercial paper program (see Note 4 in this report), and increased accrued expenses and other current liabilities, partially offset by increased trade accounts receivable and cash and cash equivalents due to an adjustment during the first quarter which impacted invoicing as well as increased prepaid expenses and other current assets largely related to a cost report receivable from Medicare and Medicaid. Our ratio of current assets to current liabilities was 1.48 and 1.63 at June 30, 2016 and December 31, 2015, respectively.

We intend to continue to address our current cash and financing requirements using cash provided by operating activities, our existing and future credit agreements, issuances under the Commercial Paper Program (See Note 4 in this report) as well as the issuance of debt securities. In addition, when funds are required for acquisitions or to meet other needs, we expect to successfully complete long-term financing arrangements, such as the issuance of senior notes. We aim to preserve financial resources with a minimum \$500 million of committed and unutilized credit facilities.

Cash provided by operating activities depends on the collection of accounts receivable. Commercial customers and governments generally have different payment cycles. A lengthening of their payment cycles could have a material adverse effect on our capacity to generate cash flow. In addition, we could face difficulties in enforcing and collecting accounts receivable under some countries' legal systems and due to the economic conditions in some countries. Accounts receivable balances, net of valuation allowances, represented Days Sales Outstanding ("DSO") of 70 at June 30, 2016, a decrease as compared to 71 at December 31, 2015.

DSO by segment is calculated by dividing the segment's accounts receivable, as converted to U.S. dollars using the average exchange rate for the period presented, less any value added tax included in the receivables, by the average daily sales for the last twelve months of that segment, as converted to U.S. dollars using the average exchange rate for the period. Receivables and sales are adjusted for amounts related to significant acquisitions made during the periods presented. The development of DSO by reporting segment is shown in the table below:

	June 30, 2016	December 31, 2015
North America days sales outstanding	54	53
EMEA days sales outstanding	105	104
Asia-Pacific days sales outstanding	103	113
Latin America days sales outstanding	147	141
FMC-AG & Co. KGaA average days sales outstanding	70	71

The DSO increase in the North America Segment is largely due to ordinary business fluctuations. An adjustment during the first quarter which impacted invoicing was largely resolved during the second quarter. The EMEA Segment's DSO increase reflects increased sales in the region coupled with fluctuations in payments of public health care organizations. The Asia-Pacific Segment's DSO decrease reflects an improvement of payment collections in China. The Latin America Segment's DSO increase reflects periodic delays in payment of public health care organizations in certain countries.

Due to the fact that a large portion of our reimbursement is provided by public health care organizations and private insurers, we expect that most of our accounts receivable will be collectible.

Net Cash Provided By (Used In) Investing Activities

We used net cash of \$655 million and \$478 million in investing activities in the six months ended June 30, 2016 and 2015, respectively.

Capital expenditures for property, plant and equipment, net of proceeds from sales of property, plant and equipment were \$498 million and \$411 million in the first six months of 2016 and 2015, respectively. In the first six months of 2016, capital expenditures were \$298 million in the North America Segment, \$112 million at Corporate, \$60 million for the EMEA Segment, \$17 million for the Asia-Pacific Segment and \$11 million for the Latin America Segment. Capital expenditures in the first six months of 2015 were \$211 million in the North America Segment, \$121 million at Corporate, \$54 million for the EMEA Segment, \$16 million for the Asia-Pacific Segment and \$9 million for the Latin America Segment. The majority of our capital expenditures were used for maintaining existing clinics, equipping new clinics, maintenance and expansion of production facilities (primarily in Germany, the North America Segment and France) and capitalization of machines provided to our customers and for Care Coordination. Capital expenditures were approximately 6% of total revenue in the first six months of 2016 as compared to 5% for the same period in 2015.

In addition to the capital expenditures discussed above, we invested approximately \$304 million cash in the first six months of 2016, \$250 million in the North America Segment, \$42 million in the EMEA Segment, \$8 million at Corporate and \$4 million in the Latin America Segment. The investment in the first six months of 2016 is primarily related to acquisitions of dialysis clinics, available for sale financial assets, acquisitions in our hospitalist and intensivist business, and a loan provided to an equity method investee in the North America

Segment as well as the acquisition of dialysis clinics in the EMEA Segment. Additionally, during the first six months of 2016, we received \$147 million from divestitures, including an approximately \$80 million repayment of unsecured loans provided to an equity method investee in 2015 and 2016 as well as approximately \$66 million related to available for sale financial assets. In the first six months of 2015, we invested approximately \$101 million cash, primarily related to the acquisition of dialysis clinics in the amount of approximately \$77 million in the North America Segment, \$15 million in the EMEA Segment, \$7 million in the Asia-Pacific Segment and \$2 million in Corporate. Additionally, during the first six months of 2015, we received \$35 million from divestitures, including a \$21 million repayment of an unsecured loan provided to an equity method investee in 2014 as well as \$9 million from the sale of our plasma collection device manufacturing business to Fresenius Kabi USA, Inc.

We anticipate capital expenditures of \$1.0 to \$1.1 billion and expect to make acquisitions of approximately \$0.75 billion in 2016. See "Report on Expected Developments" below.

Net Cash Provided By (Used In) Financing Activities

Net cash used in financing activities was \$51 million in the first six months of 2016 compared to net cash used in financing activities of \$394 million in the first six months of 2015.

In the six-month period ended June 30, 2016, cash was mainly used for the payment of dividends, repayments of long-term debt and capital lease obligations, repayments of short-term debt and distributions to noncontrolling interests, largely offset by proceeds from short-term debt. In the first six months of 2015, cash was mainly used for the payment of dividends, repayments of long-term debt and short-term debt as well as distributions to noncontrolling interests, partially offset by proceeds from short-term debt, proceeds from the exercise of stock options and proceeds from short-term debt from related parties.

On May 13, 2016, we paid a dividend with respect to 2015 of \in 0.80 per share (for 2014 paid in 2015 \in 0.78). The total dividend payment was \in 244 million (\$277 million) as compared with \in 237 million (\$263 million) in the prior year.

Non-U.S. GAAP Measures for Presentation

Constant Currency

Changes in revenue include the impact of changes in foreign currency exchange rates. We use the non-GAAP financial measure at Constant Exchange Rates or Constant Currency in our filings to show changes in our revenue without giving effect to period-to-period currency fluctuations. Under U.S. GAAP, revenues received in local (non-U.S. dollar) currency are translated into U.S. dollars at the average exchange rate for the period presented. Once we translate the local currency revenues for the Constant Currency, we then calculate the change, as a percentage, of the current period revenues using the prior period exchange rates versus the prior period revenues. This resulting percentage is a non-GAAP measure referring to a change as a percentage at Constant Currency.

We believe that revenue growth is a key indication of how a company is progressing from period to period and that the non-GAAP financial measure Constant Currency is useful to investors, lenders, and other creditors because such information enables them to gauge the impact of currency fluctuations on a company's revenue from period to period. However,

we also believe that the usefulness of data on Constant Currency period-over-period changes is subject to limitations, particularly if the currency effects that are eliminated constitute a significant element of our revenue and significantly impact our performance. We therefore limit our use of Constant Currency period-over-period changes to a measure for the impact of currency fluctuations on the translation of local currency revenue into U.S. dollars. We do not evaluate our results and performance without considering both Constant Currency period-over-period changes in non-U.S. GAAP revenue on the one hand and changes in revenue prepared in accordance with U.S. GAAP on the other. We caution the readers of this report to follow a similar approach by considering data on Constant Currency period-over-period changes only in addition to, and not as a substitute for or superior to, changes in revenue prepared in accordance with U.S. GAAP. We present the fluctuation derived from U.S. GAAP revenue next to the fluctuation derived from non-GAAP revenue. Because the reconciliation is inherent in the disclosure, we believe that a separate reconciliation would not provide any additional benefit.

Delivered EBIT

As a result of the increase of noncontrolling interest holders in our operations, we believe a measure that is meaningful to investors is operating income less noncontrolling interests ("Delivered EBIT"). Delivered EBIT approximates the operating income attributable to the shareholders of FMC-AG & Co. KGaA. As such, we believe that operating income, or EBIT, is the closest comparable U.S. GAAP measure. Below is a table showing the reconciliation of Delivered EBIT to Operating Income for each of our reporting segments:

		Three months ended June 30			Six months ended June 30			
	2	2016		2015		2016		2015
	((in millions, unaudited)			(in millions,	ıdited)		
Delivered EBIT reconciliation								
Total	_	644	_	E 47	_	4 404	_	4 054
Operating income (EBIT) less noncontrolling interests	\$	641 (76)	\$	547 (69)	\$	1,181 (145)	\$	1,051 (124)
Delivered EBIT		565		478		1,036		927
North America								
Operating income (EBIT)		513		428		949		768
less noncontrolling interests		(74)		(66)		(140)		(119)
Delivered EBIT		439		362		809		649
Dialysis								
Operating income (EBIT)		488		391		914		716
less noncontrolling interests		(66)		(53)		(123)		(97)
Delivered EBIT		422		338		791		619
Care Coordination								
Operating income (EBIT)		25		37		35		52
less noncontrolling interests		(8)		(13)		(17)		(22)
Delivered EBIT		17		24		18		30
EMEA								
Operating income (EBIT)		139		134		269		275
less noncontrolling interests		0		(1)		(2)		(1)
Delivered EBIT		139		133		267		274
Asia-Pacific								
Operating income (EBIT)		75		67		140		152
less noncontrolling interests		(2)		(2)		(3)		(4)
Delivered EBIT		73		65		137		148
Latin America								
Operating income (EBIT)		16		16		27		34
less noncontrolling interests Delivered EBIT	-	16		16		27		34
Delivered EDII		10		10		21		34

Non-U.S. GAAP Measures

EBITDA

EBITDA (earnings before interest, tax, depreciation and amortization expenses) was approximately \$1,557 million, 18.1% of revenues for the six-month period ended June 30, 2016, and \$1,408 million, 17.3% of revenues for the same period of 2015. EBITDA is the basis for determining compliance with certain covenants contained in our Amended 2012 Credit Agreement, euro-denominated notes and the indentures relating to our senior notes. You should not consider 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. In addition, not all funds depicted by EBITDA are available for management's discretionary use. For example, a substantial portion of such funds are subject to contractual restrictions and functional requirements for debt service, to fund necessary capital expenditures and to meet other commitments from time to time as described in more detail elsewhere in this report. EBITDA, as calculated, may not be comparable to similarly titled measures reported by other companies. A reconciliation of EBITDA to cash flow provided by (used in) operating activities, which we believe to be the most directly comparable U.S. GAAP financial measure, is calculated as follows:

Reconciliation of EBITDA to net cash provided by (used in) operating activities

	For the six months ended June 30,			
	2016 2		2015	
		(in millions)		
Total EBITDA	\$	1,557	\$	1,408
Interest expense (net of interest income)		(208)		(204)
Income tax expense		(306)		(273)
Change in deferred taxes, net		(26)		(73)
Changes in operating assets and liabilities		(134)		(24)
Stock compensation expense		8		1
Other items, net		(34)		(3)
Net cash provided by (used in) operating activities	\$	857	\$	832

Cash flow measures

Our consolidated statement of cash flows indicates how we generated and used cash and cash equivalents. When used in conjunction with the other primary financial statements, it provides information that helps us evaluate the changes in our net assets and our financial structure (including our liquidity and solvency). The net cash provided by (used in) operating activities is used to assess whether our business can generate the cash required to make replacement and expansion investments. Net cash provided by (used in) operating activities is impacted by the profitability of our business and development of working capital, principally receivables. The financial key performance indicator of net cash provided by (used in) operating activities in percentage of revenue shows the percentage of our revenue that is available in terms of financial resources.

Free cash flow is the cash flow provided by (used in) operating activities after capital expenditures for property, plant and equipment but before acquisitions and investments. The key performance indicator used by management is free cash flow as a percentage of revenue. This represents the percentage of revenue that is available for acquisitions, dividends to shareholders, or the reduction of debt financing.

The following table shows the significant cash flow key performance indicators for the six months ended June 30, 2016 and 2015:

	For the six months ended June 30,				
	2016			2015	
		(in millions)			
Revenue	\$	8,626	\$	8,159	
Net cash provided by (used in) operating activities		857		832	
Capital expenditures		(506)		(418)	
Proceeds from sale of property, plant and equipment		8		7	
Capital expenditures, net	\$	(498)	\$	(411)	
Free cash flow		359		421	
Net cash provided by (used in) operating activities as a					
% of revenue		9.9%		10.2%	
Free cash flow as a % of revenue		4.2%		5.2%	

Care Coordination Business Metrics for Presentation

The measures for our North America Segment discussed below include current and future programs that we will be participating in and will be reflected in the discussion of our business within the North America Segment. Currently, only the sub-capitation, BPCI, ESCO programs and other shared savings programs are included within the Member Months and Medical Cost Under Management calculations below. In the future, there may be other programs that could be included in the following metrics. These metrics may be developed further in future periods. Note that due to the timing required by CMS to review the BPCI program data that we provide, estimates have been used in order to report these metrics in a timely manner.

Member Months Under Medical Cost Management

Member months under medical cost management is calculated by multiplying the number of members who are included in value-based reimbursement programs, such as Medicare Advantage plans or other value-based programs in the U.S., by the corresponding number of months these members participate in those programs ("Member Months"). In the aforementioned programs, we are assuming the risk of generating savings. The financial results will be recorded in earnings as our performance is determined. The membership offerings within Care Coordination are sub-capitation arrangements, MA-CSNPs, ESCO and BPCI programs as well as other shared savings programs. An increase in patient membership may indicate future earnings or losses as our performance is determined through these managed care programs.

Medical Cost Under Management

Medical cost under management represents the management of medical costs associated with our patient membership in value-based programs. For ESCO, BPCI and other shared savings programs, this is calculated by multiplying the Member Months in each program by the benchmark of expected medical cost per member per month. The sub-capitation and MA-CSNPs calculation multiplies the premium per member of the program per month by the number of Member Months associated with the plan, as noted above.

Care Coordination Patient Encounters

Care Coordination patient encounters represents the total patient encounters and procedures conducted by certain of our Care Coordination activities. Specifically, Care Coordination patient encounters is the sum of all encounters and procedures completed

during the period by Sound, MedSpring Urgent Care, Fresenius Vascular Care, and National Cardiovascular Partners as well as patients in our Rx BMM program.

Balance Sheet Structure

Total assets as of June 30, 2016 increased to \$26,553 million from \$25,365 million as compared to December 31, 2015. Current assets as a percent of total assets remained flat at 27% at June 30, 2016 as compared to December 31, 2015. The equity ratio, the ratio of our equity divided by total liabilities and shareholders' equity, remained stable at 41% at June 30, 2016 as compared to December 31, 2015.

Risk and Opportunities Report

a) Risk Report

For information regarding our risks please refer to Note 10 and 11 and the chapter "Management's Discussion and Analysis", specifically the "Forward-looking statements" and the "Financial Conditions and Results of Operations - Overview, legislation and growth" sections in this report. For additional information please see chapter 2 section "Risk and Opportunities Report" on pages 92-100 of the Annual Report 2015.

b) Opportunities Report

In comparison to the information contained within the Annual Report 2015, there have been no material changes for the second quarter of 2016. Please refer to chapter 2 section "Risk and Opportunities Report" on pages 100-103 of the Annual Report 2015.

Report on Expected Developments

Targets 2016

Below is a table showing our growth outlook for 2016:

Revenue ^{(1), (2)} Operating income ⁽³⁾ Delivered EBIT ⁽³⁾ Net income growth ^{(2), (3), (4)}	Growth 7 - 10% (at Constant Exchange Rates) Growth > revenue growth Growth > revenue growth 15 - 20%
Basic earnings per share growth ^{(2), (3), (4)}	based on development of net income
Capital Expenditures	\$1.0 - 1.1 billion
Acquisitions and investments	~ \$0.75 billion
Net cash provided by (used in) operating activities in % of	
revenue ⁽³⁾	> 10%
Free cash flow in % of revenue ⁽³⁾	> 4%
Debt/EBITDA Ratio	< 3.0
Employees ⁽⁵⁾	> 109,000
Research and development expenses	\$160 - 170 million

- (1) Net of patient service bad debt provision
- (2) Targets 2016 exclude contributions from acquisitions closed in 2015 and 2016
- (3) Targets 2016 exclude special items
- (4) Net income attributable to shareholders of FMC AG & Co. KGaA
- (5) Full-time equivalents

Subsequent Events

No significant activities have taken place since the balance sheet date June 30, 2016 that have a material impact on the key figures and business earnings presented. Currently, there are no other significant changes in the structure, management, legal form of the Company or on its personnel.

Recently Implemented Accounting Pronouncements

On February 18, 2015, FASB issued Accounting Standards Update 2015-02 ("ASU 2015-02"), Consolidation (Topic 810): Amendments to the Consolidation Analysis, which focuses on clarifying guidance related to the evaluation of various types of legal entities such as limited partnerships, limited liability corporations and certain security transactions for consolidation. The update is effective for fiscal years beginning after December 15, 2015, and for interim periods within fiscal years beginning after December 15, 2015. We have implemented ASU 2015-02. These types of legal entities are predominantly utilized in the U.S. The consolidation disclosures in Note 1 of our annual report 2016 will include amended disclosures in relation to this ASU.

On November 20, 2015, FASB issued Accounting Standards Update 2015-17 ("ASU 2015-17") *Income Taxes (Topic 740): Balance Sheet Classification of Deferred Taxes*, which focuses on reducing the complexity of classifying deferred taxes on the balance sheet. ASU 2015-17 eliminates the current requirement for organizations to present deferred tax liabilities and assets as current and noncurrent in a classified balance sheet and requires the classification of all deferred tax assets and liabilities as noncurrent. The update is effective for fiscal years and interim periods within those years beginning after

December 15, 2016. We adopted this ASU as of March 31, 2016. In accordance with ASU 2015-17, deferred taxes recorded as of December 31, 2015 within current assets and liabilities have been reclassified to noncurrent assets and liabilities in the amount of \$216 million and \$36 million, respectively. As a result of deferred tax netting, noncurrent assets and liabilities were then adjusted in the amount of \$168 million.

Recent Accounting Pronouncements Not Yet Adopted

On May 28, 2014, the FASB issued Accounting Standards Update 2014-09 ("ASU 2014-09"), Revenue from Contracts with Customers, Topic 606. Simultaneously, the IASB published its equivalent revenue standard, "IFRS 15," Revenue from Contracts with Customers. The standards are the result of a convergence project between FASB and the IASB. This update specifies how and when companies reporting under U.S. GAAP will recognize revenue as well as providing users of financial statements with more informative and relevant disclosures. ASU 2014-09 supersedes some guidance included in topic 605, Revenue Recognition, some guidance within the scope of Topic 360, Property, Plant, and Equipment, and some guidance within the scope of Topic 350, Intangibles - Goodwill and Other. This ASU applies to nearly all contracts with customers, unless those contracts are within the scope of other standards (for example, lease contracts or insurance contracts). With the issuance of Accounting Standards Update 2015-14 ("ASU 2015-14"), Revenue from Contracts with Customers (Topic 606): Deferral of the Effective Date on August 12, 2015, the effective date of ASU 2014-09 for public business entities, among others, was deferred from fiscal years and interim periods within those years beginning after December 15, 2016 to fiscal years and interim periods within those years beginning after December 15, 2017. Earlier adoption is permitted. We are currently evaluating the impact of ASU 2014-09, in conjunction with all amendments, on our Consolidated Financial Statements.

On January 5, 2016, FASB issued Accounting Standards Update 2016-01 ("ASU 2016-01") Financial Instruments -- Overall (Subtopic 825-10): Recognition and Measurement of Financial Assets and Financial Liabilities. ASU 2016-01 focuses on improving the recognition and measurement of financial instruments to provide users of financial statements with more decision-useful information. ASU 2016-01 affects the accounting treatment and disclosures related to financial instruments and equity instruments. The update is effective for fiscal years and interim periods within those years beginning after December 15, 2017. Earlier adoption is generally not permitted. We are currently evaluating the impact of ASU 2016-01 on our Consolidated Financial Statements.

On February 25, 2016, FASB issued Accounting Standards Update 2016-02 ("ASU 2016-02") Leases (Subtopic 842). ASU 2016-02 is expected to increase transparency and comparability by recognizing lease assets and lease liabilities from lessees on the balance sheet and disclosing key information about leasing arrangements in the financial statements. The lessor accounting is largely unchanged. The updates are effective for fiscal years and interim periods within those years beginning after December 15, 2018. Early applications of the amendments in these updates are permitted. We are currently evaluating the impact of ASU 2016-02 on our Consolidated Financial Statements.

On March 30, 2016, FASB issued Accounting Standards Update 2016-09 ("ASU 2016-09") Compensation- Stock Compensation (Topic 718): Improvements to Employee Share- Based Payment Accounting. ASU 2016-09 simplifies guidance with regard to income tax consequences for share-based payment transactions, classification of awards as equity or liabilities as well as cash flow impacts. The updates are effective for fiscal years and interim periods within those years beginning after December 15, 2016. Early adoption is permitted. We are currently evaluating the impact of ASU 2016-09 on our Consolidated Financial Statements.

On June 16, 2016, FASB issued Accounting Standards Update 2016-13 ("ASU 2016-13") Financial Instruments—Credit Losses (Topic 326): Measurement of Credit Losses on

Financial Instruments. ASU 2016-13 amends guidance on reporting credit losses for assets held at amortized cost basis and available for sale financial assets. For Securities and Exchange Commission filers, these updates are effective for fiscal years and interim periods within those years beginning after December 15, 2019. Early adoption is permitted as of the fiscal years beginning after December 15, 2018, including interim periods within those fiscal years. We are currently evaluating the impact of ASU 2016-13 on our Consolidated Financial Statements.

Financial statements

Consolidated statements of income

(unaudited) (in \$ THOUS, except share data)

		ended June 30, ended .				For the six ended Jui			
		2016		2015		2016	2015		
Net revenue:									
Health Care	\$	3,696,766	\$	3,453,921	\$	7,221,630	6,742,932		
Less: Patient service bad debt provision		125,731		109,123		236,255	215,730		
Net Health Care		3,571,035		3,344,798		6,985,375	6,527,202		
Dialysis Products		849,454		853,938		1,640,442	1,631,461		
		4,420,489		4,198,736		8,625,817	8,158,663		
Costs of revenue:									
Health Care		2,642,815		2,481,703		5,187,075	4,897,432		
Dialysis Products		379,144		419,766		722,563	779,914		
		3,021,959		2,901,469		5,909,638	5,677,346		
Gross profit		1,398,530		1,297,267		2,716,179	2,481,317		
Operating (income) expenses:									
Selling, general and administrative		732,147		722,618		1,490,602	1,377,534		
Research and development		38,764		34,483		76,238	65,421		
Income from equity method investees		(13,471)		(6,797)		(32,042)	(13,001)		
Operating income		641,090		546,963		1,181,381	1,051,363		
Other (income) expense:									
Interest income		(17,416)		(13,169)		(28,497)	(73,109)		
Interest expense		119,912		115,127		236,282	277,175		
Income before income taxes		538,594		445,005		973,596	847,297		
Income tax expense		168,395		135,372		306,700	273,233		
Net income		370,199		309,633		666,896	574,064		
Less: Net income attributable to noncontrolling interests		76,249		68,865		144,930	123,748		
Net income attributable to shareholders of FMC-AG & Co. KGaA	d	293,950	¢	240,768	4	F21 066 ¢	450,316		
Simicioners of Fricad & Co. NGAA	\$	293,950	7	270,700	7	521,966 \$	450,310		
Basic earnings per share	\$	0.96	\$	0.79	\$	1.71 \$	1.48		
Fully diluted earnings per share	¢	0.96	\$	0.79	\$	1.71 \$	1.48		
rany unacca carmings per snare	7	3.30	4	0.79	4	<u> </u>	1.70		

${\color{red}\textbf{Consolidated Statements of Comprehensive Income} \atop (\textit{unaudited})}$

(in thousands, except share data)

	For the thre	ee months	For the six months			
	ended Ju	une 30,	ended Ju	ne 30,		
	2016	2015	2016	2015		
Net Income	\$ 370,199	\$ 309,633	\$ 666,896 \$	574,064		
Gain (loss) related to cash flow hedges Actuarial gain (loss) on defined benefit	7,747	20,986	12,314	27,938		
pension plans Gain (loss) related to foreign currency	7,490	8,106	15,367	17,335		
translation Income tax (expense) benefit related to components of other comprehensive	10,896	18,798	115,995	(108,635)		
income	(4,901)	(8,742)	(9,266)	(14,666)		
Other comprehensive income						
(loss), net of tax	21,232	39,148	134,410	(78,028)		
Total comprehensive income Comprehensive income attributable to	\$ 391,431	\$ 348,781	\$ 801,306 \$	496,036		
noncontrolling interests	76,185	69,731	147,986	120,661		
Comprehensive income attributable to shareholders of						
FMC-AG & Co. KGaA	\$ 315,246	\$ 279,050	\$ 653,320 \$	375,375		

Consolidated Balance Sheets

(in thousands, except share data)

	June 30, 2016	December 31, 2015
Assets	(unaudited)	(audited)
Current assets:		
Cash and cash equivalents	\$ 722,735	\$ 549,500
Trade accounts receivable less allowance for doubtful		
accounts of \$557,854 in 2016 and \$465,790 in 2015	3,468,559	3,285,196
Accounts receivable from related parties	179,008	218,285
Inventories	1,384,279	1,340,751
Prepaid expenses and other current assets Total current assets	1,522,586 7,277,167	1,374,715 6,768,447
	7,277,207	0,7 00,1 1.7
Property, plant and equipment, net	3,662,973	3,425,574
Intangible assets	834,965	830,489
Goodwill	13,425,072	13,032,750
Deferred taxes	177,570	188,833
Investment in equity method investees	678,833	644,709
Other assets	496,171	474,452
Total assets	\$ 26,552,751	\$ 25,365,254
Liabilities and shareholders' equity		
Current liabilities:		
Accounts payable	\$ 574,636	'
Accounts payable to related parties	217,584	153,023
Accrued expenses and other current liabilities	2,652,666	2,503,137
Short-term debt	704,653	109,252
Short-term debt from related parties	3,331	19,052
Current portion of long-term debt and capital lease obligations Income tax payable	674,522	664,335
Total current liabilities	92,184 4,919,576	72,819 4,149,446
Total Cultent Habilities	4,919,370	4,149,440
Long-term debt and capital lease obligations, less current portion	7,702,475	7,853,487
Other liabilities	511,767	465,625
Pension liabilities	609,358	585,328
Income tax payable	161,458	162,500
Deferred taxes	595,261	
Total liabilities	14,499,895	13,840,886
Noncontrolling interests subject to put provisions and other temporary		
equity	1,223,821	1,028,368
Shareholders' equity:		
Ordinary shares, no par value, €1.00 nominal value, 392,462,972 shares authorized, 306,749,540 issued and 305,749,589 outstanding	380,435	387,162
Treasury stock, at cost	(136,976)	(505,014)
Additional paid-in capital	3,025,702	3,470,308
Retained earnings	8,115,771	7,870,981
Accumulated other comprehensive income (loss)	(1,204,941)	(1,336,295)
Total FMC-AG & Co. KGaA shareholders' equity	10,179,991	9,887,142
Noncontrolling interests not subject to put provisions	649,044	608,858
Total equity	10,829,035	10,496,000
Total liabilities and equity	\$ 26,552,751	\$ 25,365,254

Consolidated Statements of Cash Flows

(unaudited, in thousands)

		For the si ended J		
		2016		2015
Operating Activities:				
Net income	\$	666,896	\$	574,064
Adjustments to reconcile net income to net cash provided by				
operating activities: Depreciation and amortization		375,992		356,813
Change in deferred taxes, net		(25,696)		(72,560)
(Gain) loss on sale of fixed assets and investments		(5,749)		1,286
Compensation expense related to stock options		8,205		593
Investments in equity method investees, net		(28,292)		(3,533)
Changes in assets and liabilities, net of amounts from businesses acquired:		(==,===,		(=,===,
Trade accounts receivable, net		(127,204)		(168,463)
Inventories		(26,867)		(128,741)
Prepaid expenses, other current and non-current assets		(91,512)		68,165
Accounts receivable from related parties		(26,791)		25,969
Accounts receivable from related parties Accounts payable to related parties		61,826		48,724
Accounts payable to related parties Accounts payable, accrued expenses and other current and		01,020		40,724
non-current liabilities		62,995		161,979
Income tax payable		13,645		(31,846)
Net cash provided by (used in) operating activities	_	857,448		832,450
Investing Activities:		•		•
Purchases of property, plant and equipment		(506,099)		(417,751)
Proceeds from sale of property, plant and equipment		8,106		6,314
Acquisitions and investments, net of cash acquired, and		,		,
purchases of intangible assets		(304,248)		(100,591)
Proceeds from divestitures		146,904		34,432
Net cash provided by (used in) investing activities		(655,337)		(477,596)
Financing Activities:				
Proceeds from short-term debt		754,831		112,825
Repayments of short-term debt		(160,100)		(128,635)
Proceeds from short-term debt from related parties		43,185		53,001
Repayments of short-term debt from related parties		(59,366)		-
Proceeds from long-term debt and capital lease obligations		154		4,191
Repayments of long-term debt and capital lease obligations Increase (decrease) of accounts receivable securitization		(218,285)		(138,625)
program		(51,000)		14,250
Proceeds from exercise of stock options, net		20,938		53,762
Dividends paid		(277,176)		(263,244)
Distributions to noncontrolling interests		(144,772)		(123,754)
Contributions from noncontrolling interests		40,252		22,453
Net cash provided by (used in) financing activities		(51,339)		(393,776)
Effect of exchange rate changes on cash and cash equivalents		22,463		(12,774)
Cash and Cash Equivalents:		22,403		(12///4)
Net increase (decrease) in cash and cash equivalents		173,235		(51,696)
Cash and cash equivalents at beginning of period		549,500		633,855
Cash and cash equivalents at end of period	\$	722,735	\$	582,159
casii ana casii equivalents at ena oi penoa	7	122,133	P	302,133

Consolidated Statement of Shareholders' Equity

For the six months ended June 30, 2016 (unaudited) and year ended December 31, 2015 (audited) (in thousands, except share data)

			(11	i tilousullus,	CACCPL SHAIC C	iata)				
	Ordinary Shares		Treasur	y Stock	Additional		Accumulated Other	Total FMC-AG & Co. KGaA	Noncontrolling interests not	
	Number of shares	No par value	Number of shares	Amount	paid in capital	Retained earnings	comprehensive income (loss)	shareholders' equity	subject to put provisions	Total Equity
Balance at December 31, 2014 Proceeds from exercise of options	311,104,251	\$ 385,215	(7,548,951)	\$ (505,014)	\$ 3,546,075	\$ 7,104,780	\$ (1,087,743)	\$ 9,443,313	\$ 585,058	\$ 10,028,371
and related tax effects Compensation expense related to	1,758,820	1,947	-	-	87,065	-	-	89,012	-	89,012
stock options Vested subsidiary stock incentive	-	-	-	-	12,323	-	-	12,323	-	12,323
plans Dividends paid	-	-	-	-	(4,613)	- (263,244)	-	(4,613) (263,244)	-	(4,613) (263,244)
Purchase/ sale of noncontrolling interests	-	_	-	-	7 461	(203,244)	-		7,169	
Contributions from/ to noncontrolling interests	-	-	-	-	7,461	-	-	7,461	(100,852)	14,630 (100,852)
Expiration of put provisions and	-	-	-	-	-	-	-	-	, , ,	, ,
other reclassifications Changes in fair value of	-	-	-	-	-	-	-	-	(5,206)	(5,206)
noncontrolling interests subject to put provisions	-	-	-	-	(178,003)	-	-	(178,003)	-	(178,003)
Net income Other comprehensive income (loss)	-	-	-	-	-	1,029,445	(248,552)	1,029,445 (248,552)	124,577 (1,888)	1,154,022 (250,440)
Comprehensive income		-	_	-	_		_	780,893	122,689	903,582
Balance at December 31, 2015	312,863,071	\$ 387,162	(7,548,951)	\$ (505,014)	\$ 3,470,308	\$ 7,870,981	\$ (1,336,295)	\$ 9,887,142	\$ 608,858	\$ 10,496,000
Proceeds from exercise of options		•			•					
and related tax effects	435,469	490	-	-	20,835	-	-	21,325	-	21,325
Compensation expense related to										
stock options Vested subsidiary stock incentive	-	-	-	-	8,205	-	-	8,205	-	8,205
plans Withdrawal of treasury stock	- (6,549,000)	(7,217)	6,549,000	368,038	(4,351) (360,821)	-	-	(4,351)	-	(4,351)
Dividends paid	(0,549,000)	(7,217)	6,349,000	300,036	(300,821)	(277,176)	-	(277,176)	-	(277,176)
Purchase/ sale of noncontrolling										
interests Contributions from/ to	-	-	-	-	22,776	-	-	22,776	22,895	45,671
noncontrolling interests Expiration of put provisions and	-	-	-	-	-	-	-	-	(45,090)	(45,090)
other reclassifications Changes in fair value of	-	-	-	-	-	-	-	-	3,848	3,848
noncontrolling interests subject to put provisions					(131,250)			(131,250)		(131,250)
Net income	_		_	_	(131,230)	521,966		521,966	56,073	578,039
Other comprehensive income (loss)	_	_	_	_	_	521,900	131,354	131,354	2,460	133,814
Comprehensive income	_	_	_	_	_	_	131,334	653,320	58,533	711,853
•										/ 11/000
Balance at June 30, 2016	306,749,540	\$ 380,435	(999.951)	\$ (136.976)	\$ 3,025,702	\$ 8.115.771	\$ (1.204.941)	\$ 10,179,991	\$ 649,044	\$ 10,829,035

Notes to Consolidated Financial Statements

(unaudited)
(in thousands, except share and per share data)

1. The Company and Basis of Presentation

The Company

Fresenius Medical Care AG & Co. KGaA ("FMC-AG & Co. KGaA" or the "Company"), a German partnership limited by shares (Kommanditgesellschaft auf Aktien), is the world's largest kidney dialysis company. The Company provides dialysis treatment and related dialysis care services to persons who suffer from end-stage renal disease ("ESRD"), as well as other health care services. The Company provides dialysis products for the treatment of ESRD, including products manufactured and distributed by the Company such as hemodialysis machines, peritoneal cyclers, dialyzers, peritoneal solutions, hemodialysis concentrates, solutions and granulates, bloodlines, renal pharmaceuticals and systems for water treatment. The Company supplies dialysis clinics it owns, operates or manages with a broad range of products in addition to sales of dialysis products to other dialysis service providers. The Company describes its other health care services as "Care Coordination." Care Coordination currently includes the coordinated delivery of pharmacy services, vascular, cardiovascular and endovascular specialty services, nondialysis laboratory testing services, physician services, hospitalist and intensivist services, health plan services and urgent care services, which, together with dialysis care services represent the Company's health care services.

In these unaudited consolidated financial statements, "FMC-AG & Co. KGaA," or the "Company," "we," "us" or "our" refers to the Company or the Company and its subsidiaries on a consolidated basis, as the context requires. The term "North America Segment" refers to the North America operating segment; the term "EMEA Segment" refers to the Europe, Middle East and Africa operating segment, the term "Asia-Pacific Segment" refers to the Asia-Pacific operating segment, and the term "Latin America Segment" refers to the Latin America operating segment. For further discussion of the Company's operating segments, see Note 13.

Basis of Presentation

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

The consolidated financial statements at June 30, 2016 and for the three and six months ended June 30, 2016 and 2015 contained in this report are unaudited and should be read in conjunction with the consolidated financial statements contained in the Company's Annual Report 2015. 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. Such financial statements reflect all adjustments that, in the opinion of management, are necessary for a fair presentation of the results of the periods presented. All such adjustments are of a normal recurring nature.

The accounting policies applied in the accompanying consolidated financial statements are the same as those applied in the consolidated financial statements at and for the year ended December 31, 2015, contained in the Company's Annual Report 2015.

Notes to Consolidated Financial Statements

(unaudited)
(in thousands, except share and per share data)

Certain items in the prior year's comparative consolidated financial statements have been reclassified to conform to the current year's presentation. Deferred taxes which were classified as current at December 31, 2015, were reclassified to noncurrent as of January 1, 2016 in accordance with Accounting Standards Update 2015-17, *Income Taxes (Topic 740) Balance Sheet Classification of Deferred Taxes*. Deferred taxes in current assets and liabilities have been reclassified to noncurrent assets and liabilities in the amount of \$216,127 and \$36,399, respectively. As a result of deferred tax netting, noncurrent assets and liabilities have been adjusted in the amount of \$168,232.

The results of operations for the three and six months ended June 30, 2016 are not necessarily indicative of the results of operations for the year ending December 31, 2016.

2. Related Party Transactions

The Company's parent, Fresenius SE & Co. KGaA ("Fresenius SE"), a German partnership limited by shares, owns 100% of the share capital of Fresenius Medical Care Management AG, the Company's general partner ("General Partner"). Fresenius SE is also the Company's largest shareholder and owns approximately 30.9% of the Company's outstanding shares at June 30, 2016. The Company has entered into certain arrangements for services, leases and products with Fresenius SE or its subsidiaries and with certain of the Company's equity method investees as described in item a) below. The Company's terms related to the receivables or payables for these services, leases and products are generally consistent with the normal terms of the Company's ordinary course of business transactions with unrelated parties. Financing arrangements as described in item b) below have agreed upon terms which are determined at the time such financing transactions occur and reflect market rates at the time of the transaction. The relationship between the Company and its key management personnel who are considered to be related parties is described in item c) below. Our related party transactions are settled through Fresenius SE's cash management system where appropriate.

a) Service Agreements, Lease Agreements and Products

The Company is party to service agreements with Fresenius SE and certain of its affiliates (collectively the "Fresenius SE Companies") to receive services, including, but not limited to: administrative services, management information services, employee benefit administration, insurance, information technology services, tax services and treasury management services. The Company also provides central purchasing services to the Fresenius SE Companies. The Company provides certain administrative services to one of its equity method investees. In 2015, the Company also performed marketing and distribution services for certain of its equity method investees. These related party agreements generally have a duration of 1-5 years and are renegotiated on an as needed basis when the agreement comes due.

The Company is a party to real estate operating lease agreements with the Fresenius SE Companies, which include leases for the Company's corporate headquarters in Bad Homburg, Germany and production sites in Schweinfurt and St. Wendel, Germany. The majority of the leases expire in 2016 and the Company intends to extend these leases.

In addition to the above mentioned service and lease agreements, the Company sold products to the Fresenius SE Companies and made purchases from the Fresenius SE Companies and equity method investees. In addition, Fresenius Medical Care Holdings, Inc. ("FMCH") purchases heparin supplied by Fresenius Kabi USA, Inc. ("Kabi USA"),

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

through an independent group purchasing organization ("GPO"). Kabi USA is an indirect, wholly-owned subsidiary of Fresenius SE. The Company has no direct supply agreement with Kabi USA and does not submit purchase orders directly to Kabi USA. FMCH acquires heparin from Kabi USA, through the GPO contract, which was negotiated by the GPO at arm's length on behalf of all members of the GPO.

The Company entered into an agreement with a Fresenius SE company for the manufacturing of plasma collection devices. The Company agreed to produce 3,500 units which can be further increased to a maximum of 4,550 units, over the length of the five year contract. On January 1, 2015, this manufacturing business was sold to Kabi USA for \$9,327 for which a fairness opinion was obtained from a reputable global accounting firm. The disposal was accounted for as a transaction between parties under common control at the carrying amounts without the generation of profits.

In December 2010, the Company formed a renal pharmaceutical company with Galenica Ltd., named Vifor Fresenius Medical Care Renal Pharma Ltd. ("VFMCRP"), an equity method investee of which the Company owns 45%. The Company has entered into exclusive supply agreements to purchase certain pharmaceuticals from VFMCRP."

Below is a summary, including the Company's receivables from and payables to the indicated parties resulting from the above described transactions with related parties.

	S	ervi	ce Agree	me	nts, Leas	se A	Agreeme	nt	s and Produ	cts	;				
			months		For the s										
		ended June 30, 2016			ended				June 30,				December 31,		
	Sales of		2016 Irchases		June 3		rchases		201	6			201	5	
	goods		goods		oods		goods								
	and	ar		ar		an	3		Accounts	Δ	counts	Δ	ccounts	Δα	counts
	services		rvices		ervices		rvices		Receivables						yables
Service Agreements															
Fresenius SE	101		12,047		97		10,388		70		4,743		422		3,185
Fresenius SE affiliates	1,700		42,307		3,784		37,869		648		2,175		2,104		4,079
Equity method investees	8,392		_		8,021		-		616		_		10,180		-
Total	\$ 10,193	\$	54,354	\$	11,902	\$	48,257	5	1,334	\$	6,918	\$	12,706	\$	7,264
Lease Agreements															
Fresenius SE	-		5,206		-		4,741		-		-		-		-
Fresenius SE affiliates			7,595		_		7,320		-		_		-		_
Total	\$ -	\$	12,801	\$	-	\$	12,061	9	-	\$	-	\$	_	\$	
Products															
Fresenius SE	2		_		4		_		_		_		_		_
Fresenius SE affiliates	12,879		18,758		13,247		18,706		8,587		5,141		8,774		3,768
Equity method investees			182,820				54,259		-		34,583				8,253
Total	\$ 12,881	\$	201,578	\$	13,251	\$	72,965	5	\$ 8,587	\$	39,724	\$	8,774	\$	12,021

b) Financing

The Company receives short-term financing from and provides short-term financing to Fresenius SE. The Company also utilizes Fresenius SE's cash management system for the settlement of certain intercompany receivables and payables with its subsidiaries and other related parties. As of June 30, 2016 and December 31, 2015, the Company had accounts receivables from Fresenius SE related to short-term financing in the amount of \$168,416 and \$131,252, respectively. As of June 30, 2016 and December 31, 2015, the

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

Company had accounts payables to Fresenius SE related to short-term financing in the amount of \$169,005 and \$115,932, respectively. The interest rates for these cash management arrangements are set on a daily basis and are based on the then-prevailing overnight reference rate for the respective currencies.

On August 19, 2009, the Company borrowed epsilon1,500 (\$1,665 at June 30, 2016 and \$1,633 at December 31, 2015) from the General Partner on an unsecured basis at 1.335%. The loan repayment has been extended periodically and is currently due August 22, 2016 with an interest rate of 1.334%. On November 28, 2013, the Company borrowed an additional epsilon1,500 (\$1,665 at June 30, 2016 and \$1,633 at December 31, 2015) with an interest rate of 1.875% from the General Partner. This loan is due on November 25, 2016 with an interest rate of 1.223%.

The Company provided unsecured term loans to one of its equity method investees during 2015 and 2016 in the amount of CHF 78,416 (\$79,840 based upon the average exchange rate for the six months ended June 30, 2016). These loans were repaid in full during the three months ended June 30, 2016.

At June 30, 2016, the Company provided a cash advance to Fresenius SE in the amount of $\\\in 12,200$ (\$13,544 at June 30, 2016) on an unsecured basis at an interest rate of 0.800%. At December 31, 2015, the Company borrowed from Fresenius SE in the amount of in 14,500 (\$15,786 at December 31, 2015) on an unsecured basis at an interest rate of 0.970%. For further information on these loan agreements, see Note 4."

c) Key Management Personnel

Due to the legal form of a German partnership limited by shares, the General Partner holds a key management position within the Company. In addition, as key management personnel, members of the Management Board and the Supervisory Board, as well as their close relatives, are considered related parties.

The Company's Articles of Association provide that the General Partner shall be reimbursed for any and all expenses in connection with management of the Company's business, including remuneration of the members of the General Partner's supervisory board and the members of the General Partner's management board. The aggregate amount reimbursed to the General Partner was \$11,967 and \$7,519, respectively, for its management services during the six months ended June 30, 2016 and 2015. As of June 30, 2016 and December 31, 2015, the Company had accounts receivable from the General Partner in the amount of \$671 and \$486, respectively. As of June 30, 2016 and December 31, 2015, the Company had accounts payable to the General Partner in the amount of \$1,937 and \$17,806, respectively.

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

3. Inventories

At June 30, 2016 and December 31, 2015, inventories consisted of the following:

	June 30,	December 31,
	 2016	 2015
Finished goods	\$ 745,230	\$ 670,291
Health care supplies	335,867	395,342
Raw materials and purchased components	225,526	206,525
Work in process	 77,656	 68,593
Inventories	\$ 1,384,279	\$ 1,340,751

4. Short-Term Debt and Short-Term Debt from Related Parties

At June 30, 2016 and December 31, 2015, short-term debt and short-term debt from related parties consisted of the following:

			L	December
	j	lune 30,		31,
		2016		2015
Borrowings under lines of credit	\$	94,130	\$	109,230
Commercial Paper Program		610,523		-
Other financial liabilities		_		22
Short-term debt	\$	704,653	\$	109,252
Short-term debt from related parties (see Note 2.b)		3,331		19,052
Short-term debt and short-term debt from		_		_
related parties	\$	707,984	\$	128,304

The Company and certain consolidated entities operate a multi-currency notional pooling cash management system. The Company met the conditions to offset balances within this cash pool for reporting purposes. At June 30, 2016, there were no offsets under the cash management system. At December 31, 2015, cash and borrowings under lines of credit in the amount of \$48,277 were offset under this cash management system.

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

Short-term Debt from related parties

The Company is party to an unsecured loan agreement with Fresenius SE under which the Company or its subsidiaries may request and receive one or more short-term advances up to an aggregate amount of \$400,000 until maturity on October 30, 2017. The interest on the advance(s) will be at a fluctuating rate per annum equal to LIBOR or EURIBOR as applicable plus an applicable margin. Advances can be repaid and reborrowed. At June 30, 2016, there were no advances from Fresenius SE under this facility. At December 31, 2015, the Company borrowed from Fresenius SE €14,500 (\$15,786 at December 31, 2015) on an unsecured basis. For further information on short-term debt from related parties, see Note 2 b).

5. Long-term Debt and Capital Lease Obligations

As of June 30, 2016 and December 31, 2015, long-term debt and capital lease obligations consisted of the following:

	-	June 30, 2016	De	ecember 31, 2015
Amended 2012 Credit Agreement Senior Notes Equity-neutral convertible bonds Accounts Receivable Facility Capital lease obligations	\$	2,480,867 5,359,276 419,224 - 47,393	\$	2,611,580 5,325,618 407,705 50,185 40,621
Other		70,237		82,113
Long-term debt and capital lease obligations Less current portion	\$	8,376,997 (674,522)	\$	8,517,822 (664,335)
Long-term debt and capital lease obligations, less current portion	\$	7,702,475	\$	7,853,487

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(unaudited)

(in thousands, except share and per share data)

Amended 2012 Credit Agreement

The following table shows the available and outstanding amounts under the Amended 2012 Credit Agreement at June 30, 2016 and December 31, 2015:

	Maximum Am June 3		Outstanding 0, 2016 ⁽¹⁾			
Revolving Credit USD	\$ 1,000,000	\$ 1,000,000	\$ -	\$ -		
Revolving Credit EUR	€ 400,000	\$ 444,080	€ -	\$ -		
USD Term Loan	\$ 2,200,000	\$ 2,200,000	\$ 2,200,000	\$ 2,200,000		
EUR Term Loan	€ 264,000	\$ 293,093	€ 264,000	\$ 293,093		
		\$ <u>3,937,173</u>	=	\$ <u>2,493,093</u>		
	Maximum Am	nount Available	Balance C	Dutstanding		
	Decembe	r 31, 2015	December	31, 2015 ⁽¹⁾		
Revolving Credit USD	\$ 1,000,000	\$ 1,000,000	\$ 25,110	\$ 25,110		
Revolving Credit EUR	€ 400,000	\$ 435,480	€ -	\$ -		
USD Term Loan	\$ 2,300,000	\$ 2,300,000	\$ 2,300,000	\$ 2,300,000		
EUR Term Loan	€ 276,000	\$ 300,481	€ 276,000	\$ 300,481		
		\$ 4,035,961	_	\$ 2,625,591		

⁽¹⁾ Amounts shown are excluding debt issuance costs.

At June 30, 2016 and December 31, 2015, the Company had letters of credit outstanding in the amount of \$3,550 and \$3,600, respectively, under the USD revolving credit facility, which are not included above as part of the balance outstanding at those dates, but which reduce available borrowings under the applicable revolving credit facility.

Accounts Receivable Facility

The following table shows the available and outstanding amounts under the Accounts Receivable Facility at June 30, 2016 and at December 31, 2015:

	Maximum Amount Available ⁽¹⁾					Balance	Οι	Outstanding ⁽²⁾			
		lune 30,	D	ecember 31,		lune 30,		Dece	ember 31,		
		2016		2015		2016		4	2015		
Accounts Receivable Facility	\$	800,000	\$	800,000	\$	-	= :	\$	51,000		

⁽¹⁾ Subject to availability of sufficient accounts receivable meeting funding criteria.

⁽²⁾ Amounts shown are excluding debt issuance costs.

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The Company also had letters of credit outstanding under the Accounts Receivable Facility in the amount of \$13,822 and \$16,622 at June 30, 2016 and December 31, 2015, respectively. These letters of credit are not included above as part of the balance outstanding at June 30, 2016 and December 31, 2015; however, they reduce available borrowings under the Accounts Receivable Facility.

6. Earnings Per Share

The following table contains reconciliations of the numerator and denominators of the basic and diluted earnings per share computations for the three and six months ended June 30, 2016 and 2015:

		For the the ended .		For the six months ended June 30,			
		2016		2015	2016	2015	
Numerator: Net income attributable to shareholders of FMC-AG & Co. KGaA	\$	293,950	\$	240,768	\$ 521,966	\$	450,316
Denominators: Weighted average number of							
Ordinary shares outstanding Potentially dilutive Ordinary		305,507,271		304,172,400	305,416,228		303,929,089
shares		258,027		1,155,218	228,752		1,052,769
Total weighted average Ordinary shares outstanding assuming dilution	,	305,765,298		305,327,618	 305,644,980		304,981,858
Basic earnings per share Fully diluted earnings per share	\$ \$	0.96 0.96				\$ \$	1.48 1.48

By resolution of the Company's annual general meeting on May 12, 2011, the Company was authorized to conduct a share buy-back program to repurchase ordinary shares. The buy-back program commenced on May 20, 2013 and was completed on August 14, 2013 after 7,548,951 shares had been repurchased in the amount of €384,966 (\$505,014). On February 16, 2016, the Company retired 6,549,000 of the repurchased shares from the buy-back program at an average weighted price of €51 per share (\$57 per share on February 16, 2016).

7. Employee Benefit Plans

The Company currently has two principal pension plans, one for German employees, the other covering employees in the United States, the latter of which was curtailed in 2002. Plan benefits are generally based on years of service and final salary. As there is no legal requirement in Germany to fund defined benefit plans, the Company's pension obligations in Germany are unfunded. Each year FMCH contributes to the plan covering

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United States employees at least the minimum required by the Employee Retirement Income Security Act of 1974, as amended.

The following table provides the calculations of net periodic benefit cost for the three and six months ended June 30, 2016 and 2015, respectively.

	For the three months ended June 30,			For the size end June				
		2016		2015	2016	2016 20		
Components of net periodic benefit cost:								
Service cost	\$	6,859	\$	6,149	13,684	\$	12,521	
Interest cost		7,246		6,972	14,575		13,915	
Expected return on plan assets		(3,868)		(4,104)	(7,740)		(8,202)	
Amortization of unrealized losses		7,519		8,106	15,426		17,335	
Amortization of prior service cost		(29)		-	(59)		_	
Net periodic benefit costs	\$	17,727	\$	17,123	35,886	\$	35,569	

8. Noncontrolling Interests Subject to Put Provisions and Other Temporary Equity

The Company has potential obligations to purchase the noncontrolling interests held by third parties in certain of its consolidated subsidiaries. These obligations are in the form of put provisions and are exercisable at the third-party owners' discretion within specified periods as outlined in each specific put provision. If these put provisions were exercised, the Company would be required to purchase all or part of third-party owners' noncontrolling interests at the appraised fair value at the time of exercise. The methodology the Company uses to estimate the fair values of the noncontrolling interest subject to put provisions assumes the greater of net book value or a multiple of earnings, based on historical earnings, development stage of the underlying business and other factors. Additionally, there are put provisions that are valued by an external valuation firm. The external valuation estimates the fair values using a combination of discounted cash flows and a multiple of earnings and/or revenue. The estimated fair values of the noncontrolling interests subject to these put provisions can also fluctuate, the discounted cash flows and the implicit multiple of earnings and/or revenue at which these noncontrolling interest obligations may ultimately be settled could vary significantly from our current estimates depending upon market conditions.

At June 30, 2016 and December 31, 2015, the Company's potential obligations under these put options were \$1,216,237 and \$1,023,755. At June 30, 2016 and December 31, 2015, put options with an aggregate purchase obligation of \$279,876 and \$258,552, respectively, were exercisable. No put options were exercised during the first six months of 2016.

The following is a roll forward of noncontrolling interests subject to put provisions for the six months ended June 30, 2016 and the year ended December 31, 2015:

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	June 30, 2016	De	cember 31, 2015
Beginning balance as of January 1, Contributions to noncontrolling interests Purchase/ sale of noncontrolling interests Contributions from noncontrolling interests Expiration of put provisions and other reclassifications Changes in fair value of noncontrolling interests Net income Other comprehensive income (loss)	\$ 1,023,755 (81,806) 45,472 11,961 (3,848) 131,250 88,857 596	\$	824,658 (164,830) 7,915 16,749 5,206 178,003 159,127 (3,073)
Ending balance as of June 30, 2016 and December 31, 2015	\$ 1,216,237	\$	1,023,755

In addition to the amounts in the table above, Other Temporary Equity related to subsidiary stock incentive plans was \$7,584 and \$4,613 as of June 30, 2016 and December 31, 2015, respectively.

9. Sources of Revenue

Outside of the U.S., the Company does not recognize patient service revenue at the time the services are rendered without assessing the patient's ability to pay. Accordingly, the additional disclosure requirements introduced with ASU 2011-07 apply solely to U.S. patient service revenue. Below is a table showing the sources of our U.S. patient service revenue (net of contractual allowance and discounts but before patient service bad debt provision), included in the Company's Health Care revenue, for the six months ended June 30, 2016 and 2015.

	June 30, 2016	June 30, 2015
Medicare program Private/alternative payors Medicaid and other government sources Hospitals	\$ 2,582,988 2,608,736 317,420 497,416	2,336,037 264,900
Total patient service revenue	\$ 6,006,560	\$ 5,508,758

10. Commitments and Contingencies

Legal and Regulatory Matters

The Company is routinely involved in numerous claims, lawsuits, regulatory and tax audits, investigations and other legal matters arising, for the most part, in the ordinary course of its business of providing health care services and products. Legal matters that the Company currently deems to be material or noteworthy are described below. For the matters described below in which the Company believes a loss is both reasonably possible and estimable, an estimate of the loss or range of loss exposure is provided. For

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the other matters described below, the Company believes that the loss probability is remote and/or the loss or range of possible losses cannot be reasonably estimated at this time. The outcome of litigation and other legal matters is always difficult to predict accurately and outcomes that are not consistent with the Company's view of the merits can occur. The Company believes that it has valid defenses to the legal matters pending against it and is defending itself vigorously. Nevertheless, it is possible that the resolution of one or more of the legal matters currently pending or threatened could have a material adverse effect on its business, results of operations and financial condition.

Commercial Litigation

On April 5, 2013, the U.S. Judicial Panel on Multidistrict Litigation ordered that the numerous lawsuits filed in various federal courts alleging wrongful death and personal injury claims against FMCH and certain of its affiliates relating to FMCH's acid concentrate products NaturaLyte[®] and GranuFlo[®] be transferred and consolidated for pretrial management purposes into a consolidated multidistrict litigation in the United States District Court for the District of Massachusetts, styled In Re: Fresenius Granuflo/Naturalyte Dialysate Products Liability Litigation, Case No. 2013-md-02428. The Massachusetts state courts and the St. Louis City (Missouri) court subsequently established similar consolidated litigation for such cases filed in Massachusetts county courts and St. Louis City court. See, In Re: Consolidated Fresenius Cases, Case No. MICV 2013-03400-O (Massachusetts Superior Court, Middlesex County). These lawsuits allege generally that inadequate labeling and warnings for these products caused harm to patients. In addition, similar cases have been filed in other state courts. On February 17, 2016, the Company reached with a committee of plaintiffs' counsel and reported to the courts an agreement in principle for settlement of potentially all cases. The agreement in principle calls for the Company to pay \$250,000 into a settlement fund in exchange for releases of all or substantially all of the plaintiffs' claims, subject to the Company's right to void the settlement under certain conditions, including if more than 3% of all plaintiffs reject the settlement or the distribution of rejecters meet certain criteria. As subsequently amended with the courts' approval as to the applicable timetable, plaintiffs must accept or reject the settlement by September 15, 2016; the Company has until October 1, 2016 to exercise any rights to void the settlement; and payment of the settlement amount must be made in October 2016 if the settlement is confirmed. The Company's affected insurers have agreed to fund \$220,000 of the settlement fund, with a reservation of rights regarding certain coverage issues between and among the Company and its insurers. The Company has accrued a net expense of \$60,000 for consummation of the settlement, including legal fees and other anticipated costs.

Subsequent to the agreement in principle, the Company's insurers in the AIG group initiated an action for declaratory judgment in New York state court advancing various arguments for reducing the amount of their coverage obligations. The Company filed an action in Massachusetts state court seeking to compel the AIG group carriers to honor their obligations under applicable policies, including reimbursement to the Company of litigation defense costs incurred before the agreement in principle was reached. The affected carriers have confirmed that the coverage litigation does not impact their commitment to fund \$220,000 of the settlement with plaintiffs in October.

Certain of the complaints in the Granuflo[®]/Naturalyte[®] litigation named combinations of FMC-AG & Co. KGaA, FMC Management AG, Fresenius SE and Fresenius Management SE as defendants, in addition to FMCH and its domestic United States affiliates. The agreement in principle provides for dismissals and releases of claims encompassing the European defendants.

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Three institutional plaintiffs have filed complaints against FMCH or its affiliates under state deceptive practices statutes resting on certain background allegations common to the GranuFlo®/NaturaLyte® personal injury litigation, but seeking as remedy the repayment of sums paid to FMCH attributable to the Granuflo®/Naturalyte® products. These cases implicate different legal standards, theories of liability and forms of potential recovery from those in the personal injury litigation and their claims will not be extinguished by the personal injury litigation settlement described above. The three plaintiffs are the Attorneys General for the States of Louisiana and Mississippi and Blue Cross Blue Shield of Louisiana. See, State of Mississippi ex rel. Hood, v. Fresenius Medical Care Holdings, Inc., No. 14-cv-152 (Chancery Court, DeSoto County); State of Louisiana ex re. Caldwell and Louisiana Health Service & Indemnity Company v. Fresenius Medical Care Airline, 2016 Civ. 11035 (U.S.D.C. D. Mass.)

Other Litigation and Potential Exposures

On February 15, 2011, a whistleblower (relator) action under the False Claims Act against FMCH was unsealed by order of the United States District Court for the District of Massachusetts and served by the relator. The United States did not intervene initially in the case United States ex rel. Chris Drennen v. Fresenius Medical Care Holdings, Inc., 2009 Civ. 10179 (D. Mass.). The relator's complaint, which was first filed under seal in February 2009, alleged that the Company sought and received reimbursement from government payors for serum ferritin and multiple forms of hepatitis B laboratory tests that were medically unnecessary or not properly ordered by a physician. Discovery on the relator's complaint closed in May 2015. On October 2, 2015, the United States Attorney moved to intervene on the relator's complaint with respect only to certain Hepatitis B surface antigen tests performed prior to 2011, when Medicare reimbursement rules for such tests changed. FMCH opposed the government's motion to intervene, which remains undecided.

Subpoenas or search warrants were issued by federal and state law enforcement authorities under the supervision of the United States Attorneys for the Districts of Connecticut, Southern Florida, Eastern Virginia and Rhode Island to American Access Care LLC ("AAC"), which the Company acquired in October 2011, and to the Company's subsidiary, Fresenius Vascular Care, Inc., which now operates former AAC centers as well as its own original facilities. As of September 30, 2015, the Company had entered into settlements of allegations made by the United States Attorneys for Connecticut, Southern Florida, and Rhode Island under which the Company paid approximately \$8,000 in exchange for releases related to activities of American Access Care prior to the acquisition. Pursuant to the AAC acquisition agreement the prior owners are obligated to indemnify the Company for payments under these settlements, subject to certain limitations and deductibles. The three settlements apply only to actions and events occurring prior to the Company's acquisition of AAC. The United States Attorney for the Eastern District of Virginia pursued a grand jury investigation against an individual surgeon employed by the Company. As of July 15, 2016, the United States Attorney advised that the grand jury investigation was being closed without charges being asserted.

On October 6, 2015, the Office of Inspector General of the United States Department of Health and Human Services ("OIG") issued a subpoena to the Company seeking information about utilization and invoicing by Fresenius Vascular Care facilities as a whole for a period beginning after the acquisition of AAC. The Company is cooperating in the

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government's inquiry, which is being managed by the United States Attorney for the Eastern District of New York.

The Company has received communications alleging conduct in countries outside the U.S. and Germany that may violate the U.S. Foreign Corrupt Practices Act ("FCPA") or other anti-bribery laws. The Audit and Corporate Governance Committee of the Company's Supervisory Board is conducting investigations with the assistance of independent counsel. The Company voluntarily advised the U.S. Securities and Exchange Commission ("SEC") and the U.S. Department of Justice ("DOJ"). The Company's investigations and dialogue with the SEC and DOJ are ongoing. The Company is cooperating with the government investigations.

Conduct has been identified that may result in monetary penalties or other sanctions under the FCPA or other anti-bribery laws. In addition, the Company's ability to conduct business in certain jurisdictions could be negatively impacted. The Company has previously recorded a non-material accrual for an identified matter. Given the current status of the investigations and remediation activities, the Company cannot reasonably estimate the range of possible loss that may result from identified matters or from the final outcome of the investigations or remediation activities.

The Company is implementing enhancements to its anti-corruption compliance program, including internal controls related to compliance with international anti-bribery laws. The Company continues to be fully committed to FCPA and other anti-bribery law compliance.

In August 2014, FMCH received a subpoena from the United States Attorney for the District of Maryland inquiring into FMCH's contractual arrangements with hospitals and physicians, including contracts relating to the management of in-patient acute dialysis services. FMCH is cooperating in the investigation.

In July 2015, the Attorney General for Hawaii issued a civil complaint under the Hawaii False Claims Act styled Hawaii v. Liberty Dialysis – Hawaii, LLC et al., Case No. 15-1-1357-07 (Hawaii 1st Circuit) alleging that Xerox State Healthcare, LLC, M Group Consulting LLC and certain Liberty subsidiaries of FMCH conspired to overbill Hawaii Medicaid for Liberty's Epogen® administrations to Hawaii Medicaid patients during the period from 2006 through 2010, prior to the time of FMCH's acquisition of Liberty. The complaint alleges that Xerox State Healthcare LLC which acted as Hawaii's contracted administrator for its Medicaid program reimbursement operations during 2006-2010, provided incorrect and unauthorized billing guidance to Liberty and its consultant, M4 Consultants, Inc. (a subsidiary of M Group Consulting LLC until 2008, and now a subsidiary of Liberty), which Liberty relied on for purposes of its Epogen® billing to the Hawaii Medicaid program. The complaint seeks civil damages authorized under the Hawaii False Claims Act.

On August 31 and November 25, 2015, respectively, FMCH received subpoenas from the United States Attorneys for the District of Colorado and the Eastern District of New York inquiring into FMCH's participation in and management of dialysis facility joint ventures in which physicians are partners. FMCH is cooperating in the investigations.

On June 30, 2016, FMCH received a subpoena from the United States Attorney for the Northern District of Texas (Dallas) seeking information about the use and management of the pharmaceutical Velphoro®, and FMCH's interactions with DaVita Healthcare

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Partners, Inc. The Company understands that the subpoena relates to the investigation previously disclosed by DaVita, and is cooperating in the investigation.

From time to time, the Company is a party to or may be threatened with other litigation or arbitration, claims or assessments arising in the ordinary course of its business. Management regularly analyzes current information including, as applicable, the Company's defenses and insurance coverage and, as necessary, provides accruals for probable liabilities for the eventual disposition of these matters.

The Company, like other healthcare providers, conducts its operations under intense government regulation and scrutiny. It must comply with regulations which relate to or govern the safety and efficacy of medical products and supplies, the marketing and distribution of such products, the operation of manufacturing facilities, laboratories and dialysis clinics, and environmental and occupational health and safety. With respect to its development, manufacture, marketing and distribution of medical products, if such compliance is not maintained, the Company could be subject to significant adverse regulatory actions by the FDA and comparable regulatory authorities outside the U.S. These regulatory actions could include warning letters or other enforcement notices from the FDA, and/or comparable foreign regulatory authority which may require the Company to expend significant time and resources in order to implement appropriate corrective actions. If the Company does not address matters raised in warning letters or other enforcement notices to the satisfaction of the FDA and/or comparable regulatory authorities outside the U.S., these regulatory authorities could take additional actions, including product recalls, injunctions against the distribution of products or operation of manufacturing plants, civil penalties, seizures of the Company's products and/or criminal prosecution. FMCH is currently engaged in remediation efforts with respect to three pending FDA warning letters. The Company must also comply with the laws of the United States, including the federal Anti-Kickback Statute, the federal False Claims Act, the federal Stark Law and the federal Foreign Corrupt Practices Act as well as 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 the Company's interpretations or the manner in which it conducts its business. 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 whistleblower actions. By virtue of this regulatory environment, the Company's business activities and practices are subject to extensive review by regulatory authorities and private parties, and continuing audits, subpoenas, other inquiries, claims and litigation relating to the Company's compliance with applicable laws and regulations. The Company may not always be aware that an inquiry or action has begun, particularly in the case of whistleblower actions, which are initially filed under court seal.

The Company operates many facilities throughout the United States and other parts of the world. 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. The Company relies upon its management structure, regulatory and legal resources, and the effective operation of its compliance program to direct, manage and monitor the activities of these employees. On occasion, the Company may identify instances where employees or other agents deliberately, recklessly or inadvertently contravene the Company's policies or violate applicable law. The actions of such persons may subject the Company and its subsidiaries to liability under the Anti-Kickback Statute, the Stark Law, the False Claims Act and the Foreign Corrupt Practices Act, among other laws and comparable laws of other countries.

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Physicians, hospitals and other participants in the healthcare 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. The Company has been and is currently subject to these suits due to the nature of its business and expects that those types of lawsuits may continue. Although the Company maintains insurance at a level which it believes to be prudent, it cannot assure that the coverage limits will be adequate or that insurance will cover all asserted claims. A successful claim against the Company or any of its subsidiaries in excess of insurance coverage could have a material adverse effect upon it and the results of its operations. Any claims, regardless of their merit or eventual outcome, could have a material adverse effect on the Company's reputation and business.

The Company has also had claims asserted against it and has had lawsuits filed against it relating to alleged patent infringements or businesses that it has acquired or divested. These claims and suits relate both to operation of the businesses and to the acquisition and divestiture transactions. The Company has, when appropriate, asserted its own claims, and claims for indemnification. A successful claim against the Company or any of its subsidiaries could have a material adverse effect upon its business, financial condition, and the results of its operations. Any claims, regardless of their merit or eventual outcome, could have a material adverse effect on the Company's reputation and business.

The Company is also subject to ongoing and future tax audits in the U.S., Germany and other jurisdictions. With respect to other potential adjustments and disallowances of tax matters currently under review, the Company does not anticipate that an unfavorable ruling could have a material impact on its results of operations. The Company is not currently able to determine the timing of these potential additional tax payments.

Other than those individual contingent liabilities mentioned above, the current estimated amount of the Company's other known individual contingent liabilities is immaterial.

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11. Financial Instruments

Non-derivative Financial Instruments

The following table presents the carrying amounts and fair values of the Company's non-derivative financial instruments at June 30, 2016, and December 31, 2015.

		June	June 30,		December 31,			
		20	16	20)15			
	Fair							
	Value	Carrying	Fair	Carrying	Fair			
	Hierarchy	Amount	Value	Amount	Value			
Assets								
Cash and cash equivalents	1	\$ 722,735	722,735	\$ 549,500	549,500			
Accounts receivable (1)(2)	2	3,663,554	3,663,554	3,521,741	3,521,741			
Available for sale financial								
assets ⁽³⁾	1	294,353	294,353	275,770	275,770			
Liabilities								
Accounts payable ⁽¹⁾	2	792,220	792,220	780,851	780,851			
Short-term debt ⁽¹⁾	2	707,984	708,071	128,304	128,304			
Long-term debt, excluding								
Amended 2012 Credit								
Agreement, Senior Notes and	_							
convertible bonds	2	117,630	118,232	172,919	172,919			
Amended 2012 Credit	2	2 400 067	2 476 064	2 611 500	2 625 501			
Agreement	2	2,480,867	2,476,964	2,611,580	2,625,591			
Senior Notes	2	5,359,276	5,821,950	5,325,618	5,782,937			
Convertible bonds	2	419,224	552,458	407,705	546,057			
Noncontrolling interests subject								
to put provisions	3	1,223,821	1,223,821	1,028,368	1,028,368			

⁽¹⁾ Also includes amounts from related parties.

The carrying amounts in the table are included in the Consolidated Balance Sheets under the indicated captions, or in the case of long-term debt, in the captions shown in Note 5.

The significant methods and assumptions used in estimating the fair values of non-derivative financial instruments are as follows:

Cash and cash equivalents are stated at nominal value which equals the fair value.

Short-term financial instruments such as accounts receivable, accounts payable and short-term debt are valued at their carrying amounts, which are reasonable estimates of the fair value due to the relatively short period to maturity of these instruments.

The fair value of available for sale financial assets quoted in an active market is based on price quotations at the period-end date.

⁽²⁾ Includes long-term accounts receivable, which are included in "Other assets" in the Consolidated Balance Sheets

⁽³⁾ Available for sale financial assets are included in "Prepaid expenses and other current assets" and "Other assets" in the Consolidated Balance Sheets.

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The fair values of major long-term financial liabilities are calculated on the basis of market information. Instruments for which market quotes are available are measured using these quotes. The fair values of the other long-term financial liabilities are calculated at the present value of the respective future cash flows. To determine these present values, the prevailing interest rates and credit spreads for the Company as of the balance sheet date are used.

The valuation of noncontrolling interests subject to put provisions is determined using significant unobservable inputs. See Note 8 for a discussion of the Company's methodology for estimating the fair value of these noncontrolling interests subject to put obligations.

Currently, there is no indication that a decrease in the value of the Company's financing receivables is probable. Therefore, the allowances on credit losses of financing receivables are immaterial.

Derivative Financial Instruments

The Company is exposed to market risk from changes in foreign exchange rates and interest rates. In order to manage the risk of currency exchange rate and interest rate fluctuations, the Company enters into various hedging transactions by means of derivative instruments with highly rated financial institutions as authorized by the Company's General Partner. On a quarterly basis, the Company performs an assessment of its counterparty credit risk. The Company currently considers this risk to be low. The Company's policy, which has been consistently followed, is that financial derivatives be used only for the purpose of hedging foreign currency and interest rate exposure.

In certain instances, the Company enters into derivative contracts that do not qualify for hedge accounting but are utilized for economic purposes ("economic hedges"). The Company does not use financial instruments for trading purposes.

The Company established guidelines for risk assessment procedures and controls for the use of financial instruments. They include a clear segregation of duties with regard to execution on one side and administration, accounting and controlling on the other.

To reduce the credit risk arising from derivatives the Company concluded Master Netting Agreements with banks. Through such agreements, positive and negative fair values of the derivative contracts could be offset against one another if a partner becomes insolvent. This offsetting is valid for transactions where the aggregate amount of obligations owed to and receivable from are not equal. If insolvency occurs, the party which owes the larger amount is obliged to pay the other party the difference between the amounts owed in the form of one net payment.

The Company elects not to offset the fair values of derivative financial instruments subject to master netting agreements in its Consolidated Balance Sheets.

At June 30, 2016 and December 31, 2015, the Company had \$17,633 and \$24,366, respectively, of derivative financial assets subject to netting arrangements and \$26,443 and \$12,765, respectively, of derivative financial liabilities subject to netting arrangements. Offsetting these derivative financial instruments would have resulted in net assets of \$8,852 and \$16,273 as well as net liabilities of \$17,662 and \$4,672 at June 30, 2016 and December 31, 2015, respectively.

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In connection with the issuance of the equity-neutral convertible bonds in September 2014, the Company purchased share options. Any change in the Company's share price above the conversion price would be offset by a corresponding value change in the share options.

Foreign Exchange Risk Management

The Company conducts business on a global basis in various currencies, though a majority of its operations are in Germany and the United States. For financial reporting purposes, the Company has chosen the U.S. dollar as its reporting currency. Therefore, changes in the rate of exchange between the U.S. dollar and the local currencies in which the financial statements of the Company's international operations are maintained affect its results of operations and financial position as reported in its consolidated financial statements.

Additionally, individual subsidiaries are exposed to transactional risks mainly resulting from intercompany purchases between production sites and other subsidiaries with different functional currencies. This exposes the subsidiaries to fluctuations in the rate of exchange between the invoicing currencies and the currency in which their local operations are conducted. For the purpose of hedging existing and foreseeable foreign exchange transaction exposures the Company enters into foreign exchange forward contracts and, on a small scale, foreign exchange options. At June 30, 2016 and December 31, 2015, the Company had no foreign exchange options.

Changes in the fair value of the effective portion of foreign exchange forward contracts designated and qualifying as cash flow hedges of forecasted product purchases and sales are reported in Accumulated Other Comprehensive Income ("AOCI"). Additionally, in connection with intercompany loans in foreign currency, the Company uses foreign exchange swaps thus assuring that no foreign exchange risks arise from those loans, which, if they qualify for cash flow hedge accounting, are also reported in AOCI. These amounts recorded in AOCI are subsequently reclassified into earnings as a component of cost of revenues for those contracts that hedge product purchases or as an adjustment of interest income/expense for those contracts that hedge loans, in the same period in which the hedged transaction affects earnings. The notional amounts of foreign exchange contracts in place that are designated and qualify as cash flow hedges totaled \$95,688 and \$193,880 at June 30, 2016 and December 31, 2015, respectively.

The Company also enters into derivative contracts for forecasted product purchases and sales and for intercompany loans in foreign currencies which do not qualify for hedge accounting but are utilized for economic hedges as defined above. In these two cases, the change in value of the economic hedge is recorded in the income statement and usually offsets the change in value recorded in the income statement for the underlying asset or liability. The notional amounts of economic hedges that do not qualify for hedge accounting totaled \$1,368,107 and \$1,637,129 at June 30, 2016 and December 31, 2015, respectively.

Interest Rate Risk Management

The Company enters into derivatives, particularly interest rate swaps and to a certain extent, interest rate options, to protect against the risk of rising interest rates. These interest rate derivatives are designated as cash flow hedges and have been entered into in order to effectively convert payments based on variable interest rates into payments at a fixed interest rate. The euro-denominated interest rate swaps expire between 2016

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

and 2019 and have a weighted average interest rate of 0.71%. Interest payable and receivable under the swap agreements is accrued and recorded as an adjustment to interest expense.

At June 30, 2016 and December 31, 2015, the notional amount of the eurodenominated interest rate swaps in place was €364,000 and €376,000 (\$404,113 and \$409,351 at June 30, 2016 and December 31, 2015, respectively).

In addition, the Company also enters into interest rate hedges ("pre-hedges") in anticipation of future long-term debt issuance, from time to time. These pre-hedges are used to hedge interest rate exposures with regard to interest rates which are relevant for the future long-term debt issuance and which could rise until the respective debt is actually issued. These pre-hedges were settled at the issuance date of the corresponding long-term debt with the settlement amount recorded in AOCI amortized to interest expense over the life of the debt. At June 30, 2016 and December 31, 2015, the Company had \$49,927 and \$58,581, respectively, related to such settlements of pre-hedges deferred in AOCI, net of tax.

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Derivative Financial Instruments Valuation

The following table shows the carrying amounts of the Company's derivatives at June 30, 2016 and December 31, 2015.

	June 3	30, 2016	December 31, 2015			
	Assets ⁽²⁾	Liabilities ⁽²⁾	Assets ⁽²⁾	Liabilities ⁽²⁾		
Derivatives in cash flow hedging relationships (1)						
Current						
Foreign exchange contracts	1,186	(1,995)	3,114	(2,921)		
Interest rate contracts	-	(666)	-	(1,637)		
Non-current						
Foreign exchange contracts	641	-	171	(127)		
Interest rate contracts		(1,927)	_	(961)		
Total	\$ 1,827	\$ (4,588) \$	3,285	(5,646)		
Derivatives not designated as hedging instruments ⁽¹⁾						
Current						
Foreign exchange contracts	26,997	(21,750)	23,908	(7,056)		
Non-current						
Foreign exchange contracts Derivatives embedded in the convertible	-	(462)	1,062	(65)		
bonds Share options to secure the convertible	-	(107,822)	-	(115,990)		
bonds	107,822	-	115,990	_		
Total	\$ 134,819	\$ (130,034) \$	140,960	\$ (123,111)		

⁽¹⁾ At June 30, 2016 and December 31, 2015, the valuation of the Company's derivatives was determined using Significant Other Observable Inputs (Level 2) in accordance with the fair value hierarchy levels established in U.S. GAAP.

The carrying amounts for the current portion of derivatives indicated as assets in the table above are included in Prepaid expenses and other current assets in the Consolidated Balance Sheets while the current portion of those indicated as liabilities are included in Accrued expenses and other current liabilities. The non-current portions indicated as assets or liabilities are included in the Consolidated Balance Sheets in Other assets or Other liabilities, respectively.

The significant methods and assumptions used in estimating the fair values of derivative financial instruments are as follows:

The fair value of interest rate swaps is calculated by discounting the future cash flows on the basis of the market interest rates applicable for the remaining term of the contract as of the balance sheet date. To determine the fair value of foreign exchange forward contracts, the contracted forward rate is compared to the current forward rate for the remaining term of the contract as of the balance sheet date. The result is then discounted on the basis of the market interest rates prevailing at the balance sheet date

⁽²⁾ Derivative instruments are marked to market each reporting period resulting in carrying amounts being equal to fair values at the reporting date.

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for the applicable currency. The fair value of the embedded derivative of the convertible bonds is calculated using the difference between the market value of the convertible bond and the market value of an adequate straight bond discounted with the market interest rates as of the reporting date.

The Company's own credit risk is incorporated in the fair value estimation of derivatives that are liabilities. Counterparty credit risk adjustments are factored into the valuation of derivatives that are assets. The Company monitors and analyses the credit risk from derivative financial instruments on a regular basis. For the valuation of derivative financial instruments, the credit risk is considered in the fair value of every individual instrument. The default probability is based upon the Credit Default Swap Spreads of each counterparty appropriate for the duration. The calculation of the credit risk considered in the valuation is performed by multiplying the default probability appropriate for the duration with the expected discounted cash flows of the derivative financial instrument.

Amount of (Gain) or

The Effect of Derivatives on the Consolidated Financial Statements

Derivatives in Cash Flow	Amount of Gain or AOCI on (Effecti for the six mon	Deriva ve Port	atives tion)	Location of (Gain) or Loss Reclassified from AOCI in Income (Effective Portion)			Loss Reclassified from AOCI in Income (Effective Portion) for the six months ended June 30,						
Hedging Relationships	2016		2015				2016		2015				
Interest rate contracts Foreign exchange	\$ (67)	\$	10,190	Interes	st e/expense	\$	12,206	\$	14,255				
contracts	1,128		(9,075)	Costs	of Revenue		(953)		12,568				
	\$ (1,061)	\$	1,115			\$	11,253	\$	26,823				
Derivatives not Designated as	Location of (Gain) or Loss Recognized in Income on	A	mount of (Gain) o Income or for the six mont	Deriva	atives								
Hedging Instruments	Derivatives		2016		2015								
Foreign exchange contracts Foreign exchange	Selling, general and administrative expense Interest	\$	23,344	\$	(20,965)								
contracts	income/expense		1,925		5,625								
Derivatives embedded in the convertible bonds	Interest income/expense		(10,513)		49,120								

For foreign exchange derivatives at June 30, 2016, the Company expects to recognize \$1,885 of losses deferred in AOCI in earnings during the next twelve months.

10,513

25<u>,269</u> \$

(49,120)

(15.340)

Share options to secure

the convertible bonds

Interest

income/expense

The Company expects to incur additional interest expense of \$21,807 over the next twelve months which is currently deferred in AOCI. This amount reflects the projected amortization of the settlement amount of the terminated swaps and the current fair value of the additional interest payments resulting from the interest rate swaps maturing between 2016 and 2019 at June 30, 2016.

Notes to Consolidated Financial Statements

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At June 30, 2016, the Company had foreign exchange derivatives with maturities of up to 18 months and interest rate swaps with maturities of up to 40 months.

12. Other Comprehensive Income (Loss)

The changes in the components of other comprehensive income (loss) for the six months ended June 30, 2016 and 2015 are as follows:

		Gain (Loss) related to cash flow hedges		ctuarial gain (loss) on defined benefit ension plans		Gain (Loss) related to foreign- currency translation		Total, before on-controlling interests		Non- ontrolling interests		Total
Balance at December 31, 2014	\$			(282,019)	\$		\$	(1,087,743)	\$	(5.261)	\$	(1,093,004)
Other comprehensive income (loss) before reclassifications Amounts reclassified from	Ψ	183	Ψ	-	٣	(105,548)	~	(105,365)	٣	(3,087)	Ψ	(108,452)
AOCI		19,528		10,896		-		30,424		-		30,424
Other comprehensive income (loss) after reclassifications		19,711		10,896		(105,548)		(74,941)		(3,087)		(78,028)
Balance at June 30, 2015	\$	(83,566)	\$	(271,123)	\$	(807,995)	\$	(1,162,684)	\$	(8,348)	\$	(1,171,032)
Balance at December 31, 2015	\$	(60,214)	\$	(225,091)	\$	(1,050,990)	\$	(1,336,295)	\$	(10,222)	\$	(1,346,517)
Other comprehensive income (loss) before reclassifications Amounts reclassified from		697		-		112,939		113,636		3,056		116,692
AOCI		8,037		9,681		-		17,718		-		17,718
Other comprehensive income (loss) after reclassifications		8,734		9,681		112,939		131,354		3,056		134,410
Balance at June 30, 2016	\$	(51,480)	\$	(215,410)	\$	(938,051)	\$	(1,204,941)	\$	(7,166)	\$	(1,212,107)

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

Reclassifications out of AOCI for the six months ended June 30, 2016 and 2015 are as follows:

Details about AOCI Components	Amo	ount of (Gain) from AOCI		Location of (Gain) Loss reclassified from AOCI in Income
		2016	 2015	
(Gain) Loss related to cash flow hedges				
Interest rate contracts	\$	12,206	\$ 14,255	Interest income/expense
Foreign exchange contracts		(953)	 12,568	Costs of Revenue
		11,253	 26,823	Total before tax
		(3,216)	 (7,295)	Tax expense or benefit
	\$	8,037	\$ 19,528	Net of tax
Actuarial (Gain) Loss on defined benefit pension plans				
Amortization of unrealized (gain)				(1)
loss		15,367	 17,335	
		15,367	 17,335	Total before tax
		(5,686)	 (6,439)	Tax expense or benefit
	\$	9,681	\$ 10,896	Net of tax
Total reclassifications for the period	\$	17,718	\$ 30,424	Net of tax

⁽¹⁾ Included in the computation of net periodic pension cost (see Note 7 for additional details).

Notes to Consolidated Financial Statements

(unaudited)
(in thousands, except share and per share data)

13. Segment and Corporate Information

The Company's operating segments are the North America Segment, the EMEA Segment, the Asia-Pacific Segment and the Latin America Segment. Management evaluates each segment using measures that reflect all of the segment's controllable revenues and expenses. With respect to the performance of business operations, management believes that the most appropriate U.S. GAAP measures are revenue, operating income and operating income margin. The Company does not include income taxes as it believes this is outside the segments' control. Financing is a corporate function, which the Company's segments do not control. Therefore, the Company does not include interest expense relating to financing as a segment measurement. Similarly, the Company does not allocate certain costs, which relate primarily to certain headquarter overhead charges, including accounting and finance, because the Company believes that these costs are also not within the control of the individual segments. Production of products, production asset management, quality management and procurement related to production are centrally managed at Corporate. The Company's global research and development is also centrally managed at Corporate. These Corporate activities do not fulfill the definition of a segment. Products are transferred to the segments at cost; therefore no internal profit is generated. The associated internal revenues for the product transfers and their elimination are recorded as Corporate activities. Capital expenditures for production are based on the expected demand of the segments and consolidated profitability considerations. In addition, certain revenues, investments and intangible assets, as well as any related expenses, are not allocated to a segment but are accounted for as Corporate.

Information pertaining to the Company's segment and Corporate activities for the three and six months ended June 30, 2016 and 2015 is set forth below.

Notes to Consolidated Financial Statements

(unaudited)

(in thousands, except share and per share data)

	North America Segment	EMEA Segment	Asia-Pacific Segment	Latin America Segment	Segment Total	Corporate	Total
Three months ended June 30, 2016							
Revenue external customers	\$ 3,167,723	\$ 676,337	\$ 396,757	\$ 174,966	\$ 4,415,783	\$ 4,706	\$ 4,420,489
Inter - segment revenue	1,116	-	4	66	1,186	(1,186)	- 4 420 400
Revenue	3,168,839	676,337	396,761	175,032	4,416,969	3,520	4,420,489
Operating income Depreciation and amortization	512,870 (107,889)	(30,958)	74,888 (12,052)	16,264 (4,248)	743,521 (155,147)	(102,431) (39,062)	(194,209)
Income (loss) from equity method investees	14,406	(711)	(785)	561	13,471	- (39,002)	13,471
Capital expenditures, acquisitions and investments	306,069	74,727	9,284	11,027	401,107	68,004	469,111
Three months ended June 30, 2015							
Revenue external customers	\$ 2,945,584	\$ 667,657	\$ 376,099	\$ 202,693	\$ 4,192,033	\$ 6,703	\$ 4,198,736
Inter - segment revenue	1,536	-	18	145	1,699	(1,699)	
Revenue	2,947,120	667,657	376,117	202,838	4,193,732	5,004	4,198,736
Operating income	428,233	134,242	67,034	15,711	645,220	(98,257)	546,963
Depreciation and amortization Income (loss) from equity method	(100,879)	(28,816)	(11,604)	(4,236)	(145,535)	(35,424)	(180,959)
investees	4,005	1,818	834	140	6,797	-	6,797
Capital expenditures, acquisitions and investments	166,171	41,434	11,006	5,442	224,053	71,197	295,250
Six months ended June 30, 2016							
Net revenue external customers	\$ 6,211,512	\$ 1,307,121	\$ 771,091	\$ 328,218	\$ 8,617,942	\$ 7,875	\$ 8,625,817
Inter - segment revenue	2,136	-	10	98	2,244	(2,244)	
Revenue	6,213,648	1,307,121	771,101	328,316	8,620,186	5,631	8,625,817
Operating Income	949,316	269,343	139,968	27,145	1,385,772	(204,391)	1,181,381
Depreciation and amortization Income (loss) from equity method	(209,214)	(59,180)	(23,599)	(7,848)	(299,841)	(76,151)	(375,992)
investees	30,939	659	(227)	671	32,042	-	32,042
Total assets	17,853,135	3,621,133	1,817,984	706,713	23,998,965	2,553,786	26,552,751
thereof investments in equity method investees	314,621	226,155	111,083	26,974	678,833	-	678,833
Capital expenditures, acquisitions and investments ⁽¹⁾	549,539	103,770	17,855	15,818	686,982	123,365	810,347
Six months ended June 30, 2015							
Net revenue external customers	\$ 5,717,063	\$ 1,296,663	\$ 729,137	\$ 400,572	\$ 8,143,435	\$ 15,228	\$ 8,158,663
Inter - segment revenue	2,826	-	18	244	3,088	(3,088)	-
Revenue	5,719,889	1,296,663	729,155	400,816	8,146,523	12,140	8,158,663
Operating Income	768,317	275,498	151,546	33,568	1,228,929	(177,566)	1,051,363
Depreciation and amortization	(198,069)	(57,142)	(22,435)	(9,049)	(286,695)	(70,118)	(356,813)
Income (loss) from equity method	0.511	2 221	1 105	44.0	10.001		10.001
investees (3) (3)	8,511	2,881	1,196	413	13,001	-	13,001
Total assets ^{(2),(3)}	16,834,078	3,463,556	1,793,006	702,015	22,792,655	2,370,226	25,162,881
thereof investments in equity method investees	280,427	221,172	111,052	25,633	638,284	-	638,284
Capital expenditures, acquisitions and investments $^{(4)}$	287,403	72,184	23,935	10,901	394,423	123,919	518,342

⁽¹⁾ North America, EMEA and Latin America acquisitions exclude \$8,370, \$90,445 and \$4,122 respectively of non-cash acquisitions for 2016.

⁽²⁾ At June 30, 2015 debt issuance costs in the amount of \$56,755 have been reclassified from Prepaid expenses and other current assets and Other assets to Long-term debt and capital lease obligations to conform to the current year's presentation.

(3) Deferred taxes which were classified as current at June 30, 2015 have been reclassified to noncurrent in accordance with Accounting Standards Update 2015-17, Income Taxes (Topic 740) Balance Sheet Classification of Deferred Taxes. Deferred taxes previously recorded in 2015 within current assets and liabilities have been reclassified to noncurrent assets and liabilities in the amount of \$223,648 and \$36,402, respectively. As a result of deferred tax netting, noncurrent assets and liabilities were then adjusted in the amount of \$190,585.

⁽⁴⁾ EMEA, Asia-Pacific and Latin America acquisitions exclude \$16,105, \$36,443 and \$250, respectively, of non-cash acquisitions for 2015.

14. Supplementary Cash Flow Information

The following additional information is provided with respect to the Consolidated Statements of Cash Flows:

	For the six months e June 30,			
		2016		2015
Supplementary cash flow information:				
Cash paid for interest	\$	194,824	\$	188,963
Cash paid for income taxes (1)	\$	330,334	\$	235,696
Cash inflow for income taxes from stock option exercises ⁽²⁾	\$	4,821	\$	11,783
Supplemental disclosures of cash flow information:				
Details for acquisitions:				
Assets acquired	\$	(403,012)	\$	(138,683)
Liabilities assumed		61,387		11,680
Noncontrolling interest subject to put provisions		43,538		15,680
Noncontrolling interest		14,112		(6,353)
Non-cash consideration		72,953		50,404
Cash paid		(211,022)		(67,272)
Less cash acquired		14,715		2,968
Net cash paid for acquisitions		(196,307)		(64,304)
Cash paid for investments		(102,905)		(14,450)
Cash paid for intangible assets		(5,036)		(21,837)
Total cash paid for acquisitions and investments, net of cash				
acquired, and purchases of intangible assets	\$	(304,248)	\$	(100,591)

⁽¹⁾ Net of tax refund.

15. Events Occurring after the Balance Sheet Date

No significant activities have taken place since the balance sheet date June 30, 2016 that have a material impact on the key figures and business earnings presented. Currently, there are no other significant changes in the structure, management, legal form of the Company or on its personnel.

⁽²⁾ Thereof the excess tax benefit allocated to additional paid-in capital for the six months ended June 30, 2016 and 2015 was \$3,829 and \$9,188, respectively.

Corporate Governance

The personally liable shareholder, represented by the Managing Board of Fresenius Medical Care Management AG, and the Supervisory Board of FMC-AG & Co. KGaA have issued a compliance declaration pursuant to 161 of the German Stock Corporation Act (AktG). The Company has frequently made this declaration available to the public by pushing it on its website: www.freseniusmedicalcare.com.

Responsibility Statement

"To the best of our knowledge, and in accordance with the applicable reporting principles for interim financial reporting, the interim consolidated financial statements give a true and fair view of the results of operations, financial position and net assets of the Fresenius Medical Care-Group, and the interim management report of the group includes a fair review of the development and performance of the business and the position of the group, together with a description of the principal opportunities and risks associated with the expected development of the group for the remaining months of the financial year."

Hof an der Saale,	July 28, 2016								
Fresenius Medical Care AG & Co. KGaA									
	Represented by the General Partner Fresenius Medical Care Management AG								
R. Powell	M. Brosnan	Harry de Wit	R. Kuerbitz						
O. Schermeier	K. Wanzek	D. Wehner							

Contact and Calendar

Contact

Fresenius Medical Care

61346 Bad Homburg

Germany

Tel. +49 6172 609 0

www.freseniusmedicalcare.com

Oliver Maier

Head of Investor Relations &

Corporate Communications

Tel. +49 6172 609 2525

Fax +49 6172 609 2301

E-Mail: ir@fmc-ag.com

Calendar 2016

Report on Third Quarter 2016

October 27, 2016

Subject to alterations