

AbbVie Inc.

Recommendation **BUY** ★ ★ ★ ★ ★

Price
\$95.89 [as of Nov 09, 2017 4:00 PM ET]

12-Mo. Target Price
\$111.00

Investment Style
Large-Cap Growth

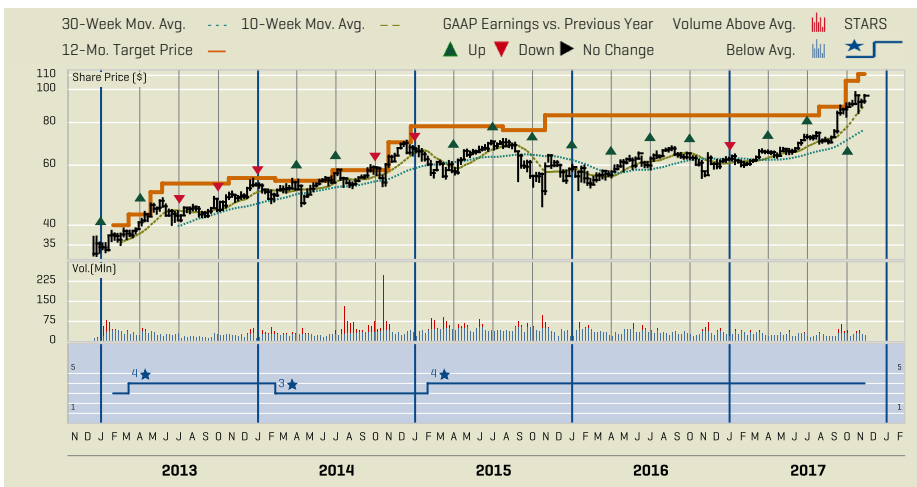
GICS Sector Health Care
Sub-Industry Biotechnology

Summary This company is a global research-based pharmaceuticals business. AbbVie's key drug is Humira, for rheumatoid arthritis and other indications.

Key Stock Statistics (Source: CFRA, S&P Global Market Intelligence (SPGMI), Company Reports)

52-Wk Range	\$98.26 - 55.87	Oper. EPS 2017E	5.54	Market Capitalization(B)	\$153.1	Beta	1.52
Trailing 12-Month EPS	4.12	Oper. EPS 2018E	6.49	Yield [%]	2.96	3-Yr Proj. EPS CAGR[%]	14
Trailing 12-Month P/E	23.25	P/E on Oper. EPS 2017E	17.28	Dividend Rate/Share	\$2.84		
\$10K Invested 5 Yrs Ago	NA	Common Shares Outstg.[M]	1,596.4	Institutional Ownership [%]	70		

Price Performance



Source: CFRA, S&P Global Market Intelligence
Analysis prepared by Equity Analyst **Jeffrey Loo** on Nov 08, 2017 06:08 PM, when the stock traded at **\$95.71**.

Highlights

- ▶ We see 2017 sales growing 9.5%, to \$28.0B. We see Humira sales of \$18.25B in 2017, up from \$16.0B in 2016, aided by greater penetration of emerging markets. We see Imbruvica [co-promoted with Johnson & Johnson] sales of \$3.1B, up from \$2.2B in 2016, but see continued decline in Viekira Pak sales to \$900M in 2017, from \$1.5B in 2016, amid pricing pressures. Yet, in August 2017, the FDA approved Mavyret to treat all six genotypes of Hepatitis C in just eight weeks. ABBV priced Mavyret at \$26,400 per treatment, below competing therapies. In September 2017, ABBV filed an NDA for Elagolix to treat endometriosis. Elagolix is also in Phase III testing for uterine fibroids.
- ▶ In September 2017, ABBV entered into a non-exclusive licensing deal with Amgen regarding Humira. The license period will begin on January 31, 2023, in the U.S. and October 16, 2018, in the EU. All Humira litigation between the companies will be dismissed. AMGN will pay ABBV royalties. In June 2016, ABBV acquired Stemcentrx for \$5.8B plus \$4B in earnouts. This follows the May 2015, \$21B acquisition of Pharmacyclics, whose Imbruvica is approved to treat three types of blood cancers.
- ▶ We see 2017 EPS of \$5.54.

Investment Rationale/Risk

- ▶ We think the shares are attractively valued, trading at 14.6X our forward 12-month EPS estimate of \$6.17, below peers. ABBV outlined its long-term objectives and sees sales rising to \$37 billion in 2020, with double-digit EPS growth through 2020. ABBV plans to launch more than 20 new products or indications through 2020. ABBV thinks operating margin could exceed 50% in 2020. We think these goals are impressive, but aggressive, particularly its new product launches. Although Humira's patent expired in December 2016, ABBV is confident that all of its other patents surrounding Humira [numbering up to 70 patents, including formulation, manufacturing, and method of treatment] will be able to protect it against biosimilar challenges at least through 2022. We believe the recent Humira licensing deal with Amgen removes an overhang on the shares. Amgen's biosimilar, Amjevita, was approved by the FDA in 2016.
- ▶ Risks include pipeline failures or if ABBV faces additional patent litigation surrounding Humira.
- ▶ Our 12-month target is \$111 based on an in-line with peers 18.0X our forward 12-month EPS estimate.

Qualitative Risk Assessment

LOW	MEDIUM	HIGH
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ABBV is heavily reliant on one drug, Humira, which currently accounts for over 60% of sales. However, recently approved drugs should help diversify sales.

Revenue/Earnings Data

Revenue (Million U.S. \$)

	1Q	2Q	3Q	4Q	Year
2018	--	--	--	--	--
2017	6,538	6,944	6,995	--	--
2016	5,958	6,452	6,432	6,796	25,638
2015	5,040	5,475	5,944	6,400	22,859
2014	4,563	4,926	5,019	5,452	19,960
2013	4,329	4,692	4,658	5,111	18,790

Earnings Per Share (U.S. \$)

	1Q	2Q	3Q	4Q	Year
2018	E 1.46	E 1.60	E 1.65	E 1.78	E 6.49
2017	1.06	1.19	1.01	E 1.46	E 5.54
2016	0.83	0.98	0.97	0.85	3.63
2015	0.64	0.83	0.74	0.92	3.13
2014	0.61	0.68	0.31	-0.51	1.10
2013	0.60	0.66	0.60	0.69	2.56

Fiscal year ended Dec 31. Next earnings report expected: Late Jan. EPS Estimates based on CFRA's Operating Earnings; historical GAAP earnings are as reported in Company reports.

Dividend Data

Amount (\$)	Date Decl.	Ex-Div. Date	Stk. of Record	Payment Date
0.64	Oct 28	Jan 11	Jan 13	Feb 15 '17
0.64	Feb 16	Apr 11	Apr 13	May 15 '17
0.64	Jun 22	Jul 12	Jul 14	Aug 15 '17
0.64	Sep 08	Oct 12	Oct 13	Nov 15 '17
0.71	Oct 27	Jan 11	Jan 12	Feb 15 '18

Business Summary November 08, 2017

CORPORATE OVERVIEW. AbbVie Inc. is a global research-based drug business that emerged as a separate company following its spin-off from Abbott Laboratories to Abbott shareholders on a share-for-share basis on January 1, 2013.

AbbVie's key product is Humira, an injectable biologic TNF [tumor necrosis factor] blocker treatment for rheumatoid arthritis (RA) and similar conditions. We estimate that Humira accounts for more than half of the global prescription drug market for rheumatoid arthritis. Besides moderate to severe RA in adults, Humira is also approved for many other uses, including juvenile idiopathic arthritis, psoriasis, ankylosing spondylitis, ulcerative colitis, Crohn's disease, and axial spondyloarthritis.

Humira's U.S. composition of matter patent expired at the end of 2016, with its equivalent European Union patent set to expire in most EU countries in April 2018. Competitors in the rheumatoid arthritis market include Remicade [marketed by Johnson & Johnson], Simponi [Johnson & Johnson], and Enbrel [Pfizer].

CORPORATE STRATEGY. AbbVie's strategic objectives include expanding Humira's sales through greater penetration of emerging markets, increased emphasis on earlier diagnosis of autoimmune patients, and new indications. ABBV also plans to launch five significant new products over the 2013-2016 time frame.

PIPELINE. AbbVie has an R&D pipeline of some 20 compounds or indications in Phase II or Phase III development across a fairly wide spectrum, including immunology, renal care, hepatitis C, women's health, oncology, multiple sclerosis, as well as Parkinson's and Alzheimer's diseases. In December 2014, the FDA approved Viekira Pak, to treat hepatitis C. Key planned launches include a levodopa-carbidopa intestinal gel [LCIG] in the U.S. for advanced Parkinson's disease; elotuzumab, a humanized monoclonal antibody for the treatment of multiple myeloma; daclizumab, a monoclonal antibody for the treatment of multiple sclerosis; ABT-199, a next-generation bcl-2 inhibitor in development for chronic lymphocytic leukemia; and new indications for Humira.

Key Phase III clinical programs include atrasentan for diabetic kidney disease and ABT-199 in chronic lymphocytic leukemia [CLL]. Other important programs are planned Phase IIB starts for: elagolix in uterine fibroids; work on a partnered JAK1 inhibitor for rheumatoid arthritis [RA]; BT-061 for RA; and ABT-719 for acute kidney injury associated with major cardiac and other surgeries. ABBV also intends to present clinical data on key development programs, including its rapidly advancing HCV program, oncology, renal disease, immunology, and Alzheimer's disease.

Corporate Information
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Chrmn & CEO

R.A. Gonzalez

EVP & CFO

W.J. Chase

COO

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EVP, Secy & General Counsel

L.J. Schumacher

EVP & CSO

M.E. Severino

Board Members

R.J. Alpern

R.S. Austin

W.H. Burnside

R.A. Gonzalez

B.J. Hart

E.M. Liddy

M.B. Meyer

E.J. Rapp

G.F. Tilton

F.H. Waddell

Domicile

Delaware

Founded

2012

Employees

29,000

Stockholders

52,270

AbbVie Inc.

Quantitative Evaluations						
Fair Value Rank	4					
	<table border="1"> <tr> <td>1</td> <td>2</td> <td>3</td> <td>4</td> <td>5</td> </tr> </table>	1	2	3	4	5
1	2	3	4	5		
	LOWEST HIGHEST Based on CFRA's proprietary quantitative model, stocks are ranked from most overvalued [1] to most undervalued [5].					
Fair Value Calculation	\$97.73 Analysis of the stock's current worth, based on CFRA's proprietary quantitative model suggests that ABBV is fairly valued.					
Volatility	<table border="1"> <tr> <td>LOW</td> <td>AVERAGE</td> <td>HIGH</td> </tr> </table>	LOW	AVERAGE	HIGH		
LOW	AVERAGE	HIGH				
Technical Evaluation	BULLISH Since October, 2017, the technical indicators for ABBV have been BULLISH.					
Insider Activity	<table border="1"> <tr> <td>UNFAVORABLE</td> <td>NEUTRAL</td> <td>FAVORABLE</td> </tr> </table>	UNFAVORABLE	NEUTRAL	FAVORABLE		
UNFAVORABLE	NEUTRAL	FAVORABLE				

Expanded Ratio Analysis				
	2016	2015	2014	2013
Price/Sales	3.98	4.24	5.28	4.51
Price/EBITDA	9.33	10.01	16.76	12.25
Price/Pretax Income	12.95	14.59	44.47	15.89
P/E Ratio	17.25	18.93	59.49	20.63
Avg. Diluted Shares Outsg. [M]	1631	1637	1610	1604

Figures based on fiscal year-end price

Key Growth Rates and Averages				
Past Growth Rate [%]		1 Year	3 Years	5 Years
Sales		12.16	10.91	8.01
Net Income		15.73	12.98	11.64
Ratio Analysis [Annual Avg.]				
Net Margin [%]		NM	NM	NM
% LT Debt to Capitalization		87.95	77.71	76.87
Return on Equity [%]		NM	NM	NM

Company Financials Fiscal year ending Dec. 31

Per Share Data [U.S. \$]	2016	2015	2014	2013	2012	2011	2010	2009	2008	2007
Tangible Book Value	-24.91	-17.97	-3.54	-2.32	-3.23	1.85	NA	NA	NA	NA
Free Cash Flow	4.05	4.31	1.84	3.63	3.81	3.74	2.87	NA	NA	NA
Earnings	3.63	3.13	1.10	2.56	3.34	2.18	2.65	NA	NA	NA
Earnings [Normalized]	3.25	3.05	1.73	2.22	2.59	2.42	2.12	NA	NA	NA
Dividends	2.35	2.10	1.75	1.60	NA	NA	NA	NA	NA	NA
Payout Ratio [%]	62	64	150	50	NA	NA	NA	NA	NA	NA
P/E Ratio: High	18.8	22.9	64.3	21.4	11.1	NA	NA	NA	NA	NA
P/E Ratio: Low	14.0	14.5	41.4	13.0	9.7	NA	NA	NA	NA	NA

Income Statement Analysis [Million U.S. \$]	2016	2015	2014	2013	2012	2011	2010	2009	2008	2007
Revenue	25,638	22,859	19,960	18,790	18,380	17,444	15,638	14,214	NA	NA
Operating Income	9,763	8,856	5,500	6,017	6,622	6,002	5,218	4,924	NA	NA
Depreciation + Amortization	1,189	836	786	897	1,150	1,272	1,184	697	NA	NA
Interest Expense	1,047	719	429	299	104	NA	NA	NA	NA	NA
Pretax Income	7,884	6,645	2,369	5,332	5,725	3,668	4,836	5,950	NA	NA
Effective Tax Rate	24.5	22.6	25.1	22.6	7.9	6.4	13.6	22.1	NA	NA
Net Income	5,953	5,144	1,774	4,128	5,275	3,433	4,178	4,637	NA	NA
Net Income [Normalized]	5,307	5,000	2,786	3,553	4,081	3,816	3,336	3,344	NA	NA

Balance Sheet and Other Financial Data [Million U.S. \$]	2016	2015	2014	2013	2012	2011	2010	2009	2008	2007
Cash	6,423	8,407	8,374	9,895	7,976	653	11	NA	NA	NA
Current Assets	16,187	16,314	16,081	17,848	15,354	7,354	8,218	NA	NA	NA
Total Assets	66,099	53,050	27,513	29,198	27,008	19,521	21,135	NA	NA	NA
Current Liabilities	9,781	10,894	11,393	6,879	6,776	5,897	3,761	NA	NA	NA
Long Term Debt	36,778	29,321	10,538	14,292	14,630	32	NA	NA	NA	NA
Total Capital	41,816	35,697	16,719	19,215	19,035	11,980	15,703	NA	NA	NA
Capital Expenditures	479	532	612	491	333	356	448	313	NA	NA
Cash from Operations	7,041	7,535	3,549	6,267	6,345	6,247	4,976	5,367	NA	NA
Current Ratio	1.65	1.50	1.41	2.59	2.27	1.25	2.19	NA	NA	NA
% Long Term Debt of Capitalization	88.0	82.1	63.0	74.4	76.9	0.3	NA	NA	NA	NA
% Net Income of Revenue	23.2	22.5	8.9	22.0	28.7	19.7	26.7	32.6	NA	NA
% Return on Assets	10.2	13.7	12.1	13.4	17.8	18.5	0.2	NA	NA	NA
% Return on Equity	NM	NM	56.9	NM	69.0	24.8	NA	NA	NA	NA

Source: S&P Global Market Intelligence. Data may be preliminary or restated; before results of discontinued operations/special items. Per share data adjusted for stock dividends; EPS diluted. E-Estimated. NA-Not Available. NM-Not Meaningful. NR-Not Ranked. UR-Under Review.

AbbVie Inc.

Sub-Industry Outlook

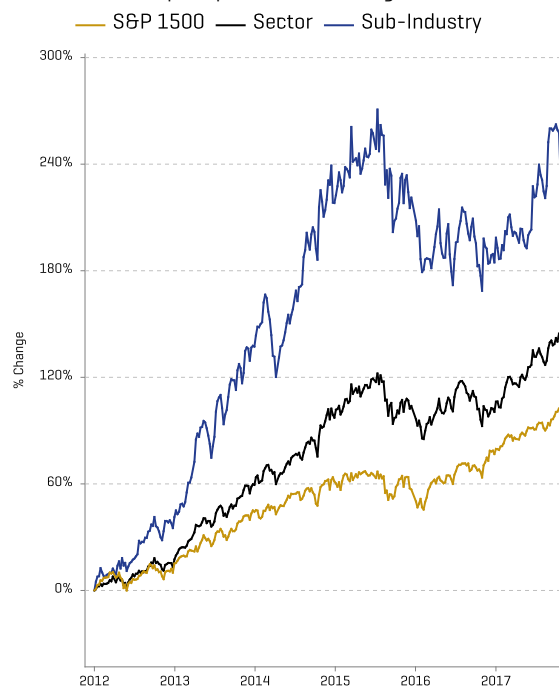
Our positive fundamental outlook for the biotechnology sub-industry for the next 12 months reflects favorable prospects for new and novel therapies to reach commercialization. In 2017, we anticipate limited sales growth of about only 0.1%, mainly due to our forecast of a 19.3% sales decline for Gilead Sciences. In 2016, biotech sales rose 7.1%. This follows the robust growth period of 2012-2015 when the compound annual growth rate was 20.2%. We think sales growth has been somewhat limited by the heightened focus on high drug prices. We are encouraged by what we view as a strong period for the reporting of late-stage clinical results, and a more accommodating U.S. FDA for approvals. However, in 2016, the FDA approved only 22 new drugs, down significantly from the robust 45 new drugs approved in 2015, the most since a record 53 were approved in 1996. We think many of these newly-approved drugs have significant commercial prospects and represent major advances in therapies for diseases such as cystic fibrosis, hepatitis C, multiple sclerosis and cancer. We expect wider adoption of biomarker research and genetic-targeted clinical studies to help bolster long-term R&D pipeline productivity. In late 2012, the FDA introduced "breakthrough therapy" designations, intended to speed development of promising programs. We expect a favorable mergers and acquisitions climate, as large pharmaceutical firms move to offset lost revenues from expiring drug patents and large biotechs bolster their drug pipelines amid maturing products. We also see large cap biotechs generating cash flows supporting larger scale acquisitions of their own. In 2011, industry bellwether Amgen became the first biotech company to initiate a regular dividend. Gilead Sciences began dividend payments in 2015. The 2010 health care reform law established the FDA's authorization to govern

"biosimilar" drug approvals and set a 12-year exclusivity to branded drug makers. We see biosimilars advancing at a modest rate over the next several years. In March 2015, the FDA approved Novartis' filgrastim biosimilar, of Amgen's Neupogen. Novartis' subsidiary, Sandoz, began selling the biosimilar under the name Zarxio in September 2015. In April 2016, the FDA approved Inflectra, manufactured by Celltrion and co-marketed with Pfizer. Inflectra is a biosimilar of Johnson and Johnson's Remicade. Pfizer began shipping Inflectra "at risk" in November 2016, at a 15% discount. We expect biosimilars to sell at more modest price discounts than generics in the pharmaceutical industry due to higher clinical, manufacturing and marketing costs, and we expect branded drugs to retain significant market share due to a lack of interchangeability. We think biosimilars may appeal to new users, but we expect current users who are stable with their current treatment would likely continue with the reference drug. But we note prescription benefit managers and health insurers are exerting more influence over drug prescriptions and pricing. In 2017, CVS removed Neupogen from its formulary and replaced it with Zarxio. In 2016, the S&P biotech index declined 14.3% vs. a 10.6% rise for the S&P 1500. Year to date through September 8, 2017, the S&P biotech index rose 26.5% vs. a 9.1% rise for the S&P 1500 Index.

Industry Performance

GICS Sector: Health Care Sub-Industry: Biotechnology

Based on S&P 1500 Indexes
Five-Year market price performance through Nov 10, 2017



NOTE: All Sector & Sub-Industry information is based on the Global Industry Classification Standard (GICS).

Past performance is not an indication of future performance and should not be relied upon as such.

Source: S&P Global Market Intelligence

Sub-Industry: Biotechnology Peer Group*: Biotechnology

Peer Group	Stock Symbol	Exchange	Currency	Recent Stock Price [\$]	Stk. Mkt. Cap. [M \$]	30-Day Price Chg. [%]	1-Year Price Chg. [%]	P/E Ratio	Yield [%]	Return on Equity [%]	LTD to Cap [%]
AbbVie Inc.	ABBV	NYSE	USD	95.89	153,082	5.2	53.1	23	3.0	NM	88.0
Actelion Ltd	ALIQ.Y	OTCPK	USD	70.02	30,061	0.3	91.7	41	5.7	52.7	NA
Alexion Pharmaceuticals, Inc.	ALXN	NasdaqGS	USD	116.37	25,998	-19.1	-8.5	52	Nil	4.7	24.1
Amgen Inc.	AMGN	NasdaqGS	USD	174.00	126,309	-6.3	18.8	16	2.6	26.6	46.8
Biogen Inc.	BIIB	NasdaqGS	USD	309.96	65,549	-6.9	-3.1	19	Nil	34.4	34.9
CSL Limited	CSLL.Y	OTCPK	USD	55.82	50,501	5.3	50.2	38	1.2	46.7	NA
Celgene Corporation	CELG	NasdaqGS	USD	102.46	80,668	-26.7	-14.7	24	Nil	31.9	66.0
Gilead Sciences, Inc.	GILD	NasdaqGS	USD	73.02	95,384	-12.1	-6.9	8	2.8	70.1	57.6
Regeneron Pharmaceuticals, Inc.	REGN	NasdaqGS	USD	406.21	42,867	-11.9	-2.9	41	Nil	22.1	NA
Shire plc	SHPG	NasdaqGS	USD	140.07	42,279	-7.5	-25.1	63	0.7	3.1	37.7
Vertex Pharmaceuticals Incorporated	VRTX	NasdaqGS	USD	148.17	37,473	-4.1	59.5	NM	Nil	-6.9	NA

*For Peer Groups with more than 10 companies or stocks, selection of issues is based on market capitalization.

NA-Not Available NM-Not Meaningful.

Note: Peers are selected based on Global Industry Classification Standards and market capitalization. The peer group list includes companies with similar characteristics, but may not include all the companies within the same industry and/or that engage in the same line of business.

AbbVie Inc.

Analyst Research Notes and other Company News

October 27, 2017

02:12 pm ET... CFRA KEEPS BUY OPINION ON SHARES OF ABBVIE INC. [ABBV 93.05****]: We raise our 12-month target \$5 to \$111 on in-line with peers 18X our forward 12-months EPS estimate of \$6.17. Q3 EPS of \$1.41 vs. \$1.21 is \$0.01 ahead of our view. We raise our '17 EPS estimate \$0.03 to \$5.54. ABBV provides '18 EPS guidance of \$6.37-\$6.57, in line with our \$6.49 view. Sales grew 9% with Humira sales up 15.8% and Imbruvica up 37.3%. It now expects Humira sales to approach \$21B in 2020, which we believe is achievable following a patent settlement with Amgen. We are also encouraged by its pipeline development with recent NDA filings for Elagolix for endometriosis. /Jeffrey Loo, CFA

September 28, 2017

12:04 pm ET... CFRA REITERATES BUY OPINION ON SHARES OF ABBVIE [ABBV 89.705****]: We raise our 12-month target \$17 to \$106 on an in-line to peers 18X our forward 12-months EPS estimate of \$5.94. ABBV announces a deal with Amgen [AMGN 187 ****] regarding Humira litigation. Under terms of the agreement, ABBV will grant AMGN a non-exclusive license to Humira's intellectual property. The license period will begin on Jan. 31, 2023, in the U.S. and Oct. 16, 2018, in the EU. All Humira litigation between the companies will be dismissed. AMGN will pay ABBV royalties. We view the deal positively for ABBV as it removes an overhang over the world's best selling drug. /Jeffrey Loo, CFA

July 28, 2017

02:15 pm ET... CFRA KEEPS BUY OPINION ON SHARES OF ABBVIE INC. [ABBV 70.21****]: We raise our 12-month target \$5 to \$89 on slightly below peers 15X our forward 12-months EPS of \$5.94. Q2 EPS of \$1.42 vs. \$1.26 is \$0.04 ahead of our estimate. We keep our '17 EPS estimate at \$5.51 but raise '18's by \$0.13 to \$6.49. Sales rose 7.6% with global Humira sales up 13.7%, and up 18.0% in the U.S., while Imbruvica sales rose 42.6%. We view ABBV's pipeline positively and await Phase III data for risankizumab in psoriasis and upadacitinib for rheumatoid arthritis. We see potential NDA filings in 2018 for both compounds. We expect data from several other trials in H2 17 /Jeffrey Loo, CFA

April 27, 2017

04:25 pm ET... CFRA KEEPS BUY OPINION ON SHARES OF ABBVIE INC. [ABBV 66.13****]: We keep our 12-month target at \$84 on slightly below peers 15.2X our 2017 EPS estimate of \$5.51. Q1 EPS of \$1.28 vs. \$1.15 is in-line with our estimate. Sales rose 9.7% on 15.1% rise in Humira sales with U.S. sales up 22.8% and international sales up 2.9%. Imbruvica sales rose 44.7% on strong uptake in first line chronic lymphocytic leukemia where we think Imbruvica is gaining market share. We look for continued robust sales of Imbruvica as we see more approvals for additional indications. ABBV also plans to submit a NDA for Elagolix to treat endometriosis later this year. /Jeffrey Loo, CFA

February 21, 2017

09:23 am ET... S&P CAPITAL IQ ADDS ABBVIE TO ITS TOTAL RETURN PORTFOLIO [ABBV 61.77****]: We see ABBV as among the fastest growing biotech companies. Although Humira's composition of matter patent expired in Dec. 2016, ABBV believes numerous other patents will be able to protect it from biosimilar competition until 2022. Trading at 11.2X our '17 EPS estimate, we think the shares are undervalued. Our 12-month target is \$84 and our EPS estimates are \$5.51 in '17 and \$6.36 in '18. ABBV pays an annual dividend of \$2.56, which has increased every year since '13. The shares yield 4.1%. ABBV replaces Gilead Sciences [GILD 70 ****] in our Total Return Model Portfolio. /C. Muir

January 27, 2017

11:53 am ET... S&P CAPITAL IQ KEEP BUY OPINION ON SHARES OF ABBVIE INC. [ABBV 59.61****]: We keep our 12-month target at \$84 on in-line to peers 15.2X our 2017 EPS estimate. We set 2018 EPS at \$6.36. Q4 EPS of \$1.20 vs. \$1.13 is \$0.01 ahead of our estimate, but sales growth of 6.9% was below our forecast partly on lower Viekira sales. However, Humira sales rose a robust 15.5%. In spite of the sales shortfall, we believe ABBV has a robust pipeline to drive future growth. We also see continued robust Humira sales as we think it will be several years before Amgen's biosimilar Amjevita reaches the market amid their patent lawsuit. ABBV guides 2017 EPS of \$5.44-\$5.54. /Jeffrey Loo, CFA

October 28, 2016

01:58 pm ET... S&P CAPITAL IQ KEEPS BUY OPINION ON SHARES OF ABBVIE INC. [ABBV 57.58****]: We keep our 12-month target at \$84 on in-line with peers 16X our forward 12-months EPS estimate of \$5.25. Q3 EPS of \$1.21 vs. \$1.13 is \$0.02 ahead of our estimate, but we lower our 2016 and 2017 EPS estimates \$0.02 and \$0.42 to \$4.81 and \$5.51. ABBV provides preliminary 2017 guidance of low double digit sales growth and EPS growth of 13%-15%. Sales rose 8% with Humira up 11%. Humira sales were slightly below our forecast on softer overseas sales, but ABBV indicated it does not believe indirect competition from Inflectra, the biosimilar for Remicade, has had an impact on sales. /Jeffrey Loo, CFA

July 29, 2016

11:50 am ET... S&P GLOBAL KEEPS BUY OPINION ON SHARES OF ABBVIE [ABBV 66.05****]: We keep our 12-month target at \$84 on in-line to peers 16.6X our forward 12-months EPS estimate of \$5.07. Q2 EPS of \$1.26, vs. \$1.08, is \$0.07 ahead of our estimate. We raise our 2016 EPS estimate \$0.07 to \$4.83. Sales rose 18% on robust growth of Humira, up 17%, and Imbruvica, driven by the approval for first line usage in chronic lymphocytic leukemia. We expect continued market share gain over the next several quarters. But Viekira sales in the U.S. is losing market share on entrance of competitor Zepatier and we see softness in EU toward year-end as Zepatier is rolled out. /Jeffrey Loo, CFA

Note: Research notes reflect S&P Global Market Intelligence's published opinions and analysis on the stock at the time the note was published. The note reflects the views of the equity analyst as of the date and time indicated in the note, and may not reflect S&P Global Market Intelligence's current view on the company.

AbbVie Inc.

Analysts' Recommendations

	No. of Recommendations	% of Total	1 Mo.Prior	3 Mos.Prior
Buy	8	38	8	7
Buy/Hold	2	10	2	2
Hold	11	52	11	12
Weak Hold	0	0	0	0
Sell	0	0	0	0
No Opinion	0	0	0	0
Total	21	100	21	21

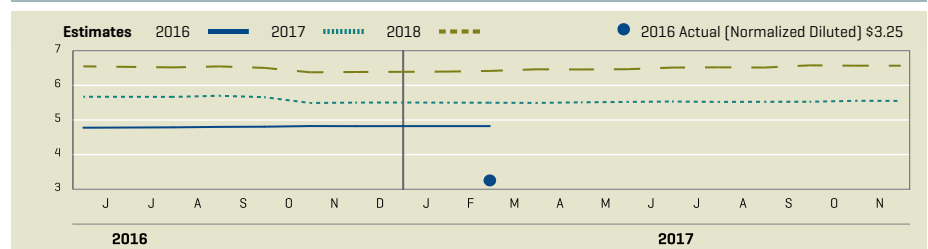
Wall Street Consensus Opinion

BUY/HOLD

Wall Street Consensus vs. Performance

For fiscal year 2017, analysts estimate that ABBV will earn USD \$5.55. For the 3rd quarter of fiscal year 2017, ABBV announced earnings per share of USD \$1.01, representing 18.2% of the total revenue estimate. For fiscal year 2018, analysts estimate that ABBV's earnings per share will grow by 18% to USD \$6.57.

Wall Street Consensus Estimates



Fiscal Years	Avg Est.	High Est.	Low Est.	# of Est.	Est. P/E
2018	6.57	6.99	6.44	18	14.6
2017	5.55	5.59	5.53	15	17.3
2018 vs. 2017	▲18%	▲25%	▲16%	▲20%	▼-16%
Q4'18	1.71	1.80	1.64	5	56.1
Q4'17	1.43	1.46	1.41	12	67.0
Q4'18 vs. Q4'17	▲20%	▲23%	▲16%	▼-58%	▼-16%

Forecasts are not reliable indicator of future performance.

Note: A company's earnings outlook plays a major part in any investment decision. S&P Global Market Intelligence organizes the earnings estimates of over 2,300 Wall Street analysts, and provides their consensus of earnings over the next two years, as well as how those earnings estimates have changed over time. Note that the information provided in relation to consensus estimates is not intended to predict actual results and should not be taken as a reliable indicator of future performance.

Note: For all tables, graphs and charts in this report that do not cite any reference or source, the source is S&P Global Market Intelligence.

AbbVie Inc.

Glossary

STARS

Since January 1, 1987, CFRA Equity and Fund Research Services, and its predecessor S&P Capital IQ Equity Research has ranked a universe of U.S. common stocks, ADRs (American Depositary Receipts), and ADSs (American Depositary Shares) based on a given equity's potential for future performance. Similarly, we have ranked Asian and European equities since June 30, 2002. Under proprietary STARS (Stock Appreciation Ranking System), equity analysts rank equities according to their individual forecast of an equity's future total return potential versus the expected total return of a relevant benchmark (e.g., a regional index [S&P Asia 50 Index, S&P Europe 350® Index or S&P 500® Index]), based on a 12-month time horizon. STARS was designed to meet the needs of investors looking to put their investment decisions in perspective. Data used to assist in determining the STARS ranking may be the result of the analyst's own models as well as internal proprietary models resulting from dynamic data inputs.

S&P Global Market Intelligence's Quality Ranking

(also known as **S&P Capital IQ Earnings & Dividend Rankings**) - Growth and stability of earnings and dividends are deemed key elements in establishing S&P Global Market Intelligence's earnings and dividend rankings for common stocks, which are designed to encapsulate the nature of this record in a single symbol. It should be noted, however, that the process also takes into consideration certain adjustments and modifications deemed desirable in establishing such rankings. The final score for each stock is measured against a scoring matrix determined by analysis of the scores of a large and representative sample of stocks. The range of scores in the array of this sample has been aligned with the following ladder of rankings:

A+	Highest	B	Below Average
A	High	B-	Lower
A-	Above Average	C	Lowest
B+	Average	D	In Reorganization
NR	Not Ranked		

EPS Estimates

CFRA's earnings per share (EPS) estimates reflect analyst projections of future EPS from continuing operations, and generally exclude various items that are viewed as special, non-recurring, or extraordinary. Also, EPS estimates reflect either forecasts of equity analysts; or, the consensus (average) EPS estimate, which are independently compiled by S&P Global Market Intelligence, a data provider to CFRA. Among the items typically excluded from EPS estimates are asset sale gains; impairment, restructuring or merger-related charges; legal and insurance settlements; in process research and development expenses; gains or losses on the extinguishment of debt; the cumulative effect of accounting changes; and earnings related to operations that have been classified by the company as discontinued. The inclusion of some items, such as stock option expense and recurring types of other charges, may vary, and depend on such factors as industry practice, analyst judgment, and the extent to which some types of data is disclosed by companies.

12-Month Target Price

The equity analyst's projection of the market price a given security will command 12 months hence, based on a combination of intrinsic, relative, and private market valuation metrics, including Fair Value.

CFRA Equity Research

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Abbreviations Used in Equity Research Reports

CAGR	- Compound Annual Growth Rate
CAPEX	- Capital Expenditures
CY	- Calendar Year
DCF	- Discounted Cash Flow
DDM	- Dividend Discount Model
EBIT	- Earnings Before Interest and Taxes
EBITDA	- Earnings Before Interest, Taxes, Depreciation & Amortization
EPS	- Earnings Per Share
EV	- Enterprise Value
FCF	- Free Cash Flow
FFO	- Funds From Operations

FY	- Fiscal Year
P/E	- Price/Earnings
P/NAV	- Price to Net Asset Value PEG Ratio - P/E-to-Growth Ratio PV - Present Value
R&D	- Research & Development ROCE - Return on Capital Employed ROE - Return on Equity
ROI	- Return on Investment
ROIC	- Return on Invested Capital
ROA	- Return on Assets
SG&A	- Selling, General & Administrative Expenses
SOTP	- Sum-of-The-Parts
WACC	- Weighted Average Cost of Capital

Dividends on American Depositary Receipts (ADRs) and American Depositary Shares (ADSs) are net of taxes (paid in the country of origin).

Qualitative Risk Assessment

Reflects an equity analyst's view of a given company's operational risk, or the risk of a firm's ability to continue as an ongoing concern. The Qualitative Risk Assessment is a relative ranking to the U.S. STARS universe, and should be reflective of risk factors related to a company's operations, as opposed to risk and volatility measures associated with share prices. For an ETF this reflects on a capitalization-weighted basis, the average qualitative risk assessment assigned to holdings of the fund.

STARS Ranking system and definition:

★★★★★ 5-STARs (Strong Buy):

Total return is expected to outperform the total return of a relevant benchmark, by a notable margin over the coming 12 months, with shares rising in price on an absolute basis.

★★★★★ 4-STARs (Buy):

Total return is expected to outperform the total return of a relevant benchmark over the coming 12 months, with shares rising in price on an absolute basis.

★★★★★ 3-STARs (Hold):

Total return is expected to closely approximate the total return of a relevant benchmark over the coming 12 months, with shares generally rising in price on an absolute basis.

★★★★★ 2-STARs (Sell):

Total return is expected to underperform the total return of a relevant benchmark over the coming 12 months, and the share price is not anticipated to show a gain.

★★★★★ 1-STAR (Strong Sell):

Total return is expected to underperform the total return of a relevant benchmark by a notable margin over the coming 12 months, with shares falling in price on an absolute basis.

Relevant benchmarks:

In North America, the relevant benchmark is the S&P 500 Index, in Europe and in Asia, the relevant benchmarks are the S&P Europe 350 Index and the S&P Asia 50 Index, respectively.

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Stocks are ranked in accordance with the following ranking methodologies:

STARS Stock Reports:

Qualitative STARS recommendations are determined and assigned by equity analysts. For reports containing STARS recommendations refer to the Glossary section of the report for detailed methodology and the definition of STARS rankings.

Quantitative Stock Reports:

Quantitative recommendations are determined by ranking a universe of common stocks based on 5 measures or model categories: Valuation, Quality, Growth, Street Sentiment, and Price Momentum. In the U.S., a sixth sub-category for Financial Health will also be displayed. Percentile scores are used to compare each company to all other companies in the same universe for each model category. The five (six) model category scores are then weighted and rolled up into a single percentile ranking for that company. For reports containing quantitative recommendations refer to the Glossary section of the report for detailed methodology and the definition of Quantitative rankings.

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STARS Stock Reports:

Global STARS Distribution as of June 30, 2017

Ranking	North America	Europe	Asia	Global
Buy	37.4%	25.9%	36.6%	35.4%
Hold	55.1%	56.1%	39.4%	53.5%
Sell	7.5%	18.0%	24.0%	11.1%
Total	100.0%	100.0%	100.0%	100.0%

Analyst Certification:

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