

1st Quarter 2026 Results

1st Quarter 2026 Sales

\$24.1B

Worldwide increased ▲
9.9%

Excluding the impact of translational currency

Stelara impacted results by ~(-540) basis points

Worldwide increased ▲
6.4%¹

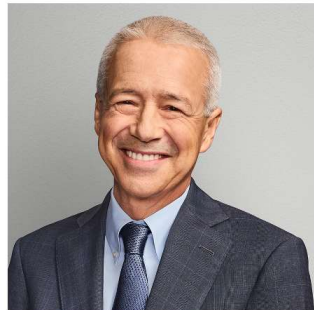
Diluted earnings per share (EPS)

\$2.14

Adjusted diluted earnings per share¹

\$2.70

Worldwide decreased ▼
(2.5)%



Joaquin Duato
Chairman & Chief Executive Officer
Johnson & Johnson

“ Johnson & Johnson had a strong start to 2026 and is delivering on its promise for a year of accelerated growth and impact. The depth and strength of our portfolio and pipeline is unrivaled and our relentless focus on innovation delivered multiple game-changing approvals this quarter, including ICOTYDE in the U.S. for moderate to severe plaque psoriasis and VARIPULSE Pro in Europe. These advancements have the potential to transform patient outcomes and create sustainable, long-term value for shareholders. ”

\$15.4 billion

Worldwide Innovative Medicine sales

Innovative Medicine worldwide reported sales increased 11.2% or 7.4% operationally². Stelara impacted results² by ~(-920) basis points. Primary operational drivers:



\$8.6 billion

Worldwide MedTech sales

MedTech worldwide reported sales increased 7.7% or 4.6% operationally². Primary operational drivers:



Electrophysiology



Abiomed



Shockwave



Trauma



Wound Closure & Healing



Contact Lenses



Biosurgery



Surgical Vision

For full financial data, non-GAAP reconciliations and cautionary statements, please refer to Johnson & Johnson's earnings release issued on April 14, 2026 available at <https://www.investor.jnj.com/financials/quarterly-results/default.aspx>

¹ Non-GAAP financial measure; non-GAAP financial measures should not be considered replacements for, and should be read together with, the most comparable GAAP financial measures.

² Non-GAAP measure; excludes the impact of translational currency.

Note: Values may be rounded

Caution Concerning Forward-Looking Statements: This document contains "forward-looking statements" as defined in the Private Securities Litigation Reform Act of 1995 regarding future operating and financial performance. You are cautioned not to rely on these forward-looking statements, which are based on current expectations of future events. For important information about the risks and uncertainties that could cause actual results to vary materially from the assumptions, expectations, and projections expressed in any forward-looking statements, review the "Note to Investors Concerning Forward-Looking Statements" included in the Johnson & Johnson earnings release issued on April 14, 2026 as well as the most recently filed Johnson & Johnson Reports on Forms 10-K and 10-Q. Johnson & Johnson does not undertake to update any forward-looking statement as a result of new information or future events or developments.

1st Quarter 2026 Earnings Call

April 14, 2026

Cautionary note on Forward-looking statements

This presentation contains “forward-looking statements” as defined in the Private Securities Litigation Reform Act of 1995 regarding, among other things: future operating and financial performance, product development, and market position and business strategy. The viewer is cautioned not to rely on these forward-looking statements. These statements are based on current expectations of future events. If underlying assumptions prove inaccurate or known or unknown risks or uncertainties materialize, actual results could vary materially from the expectations and projections of Johnson & Johnson. Risks and uncertainties include, but are not limited to: economic factors, such as interest rate and currency exchange rate fluctuations or changes to applicable laws and regulations; competition, including technological advances, new products and patents attained by competitors; challenges inherent in new product research and development, including uncertainty of clinical success and obtaining regulatory approvals; uncertainty of commercial success for new and existing products; challenges to patents; the impact of patent expirations; the ability of the Company to successfully execute strategic plans, including restructuring plans; the impact of business combinations and divestitures; manufacturing difficulties or delays, internally or within the supply chain; product efficacy or safety concerns resulting in product recalls or regulatory action; significant adverse litigation or government action, including related to product liability claims; changes to applicable laws and regulations, including tax laws and global health care reforms; trends toward health care cost containment; changes in behavior and spending patterns of purchasers of health care products and services; financial instability of international economies and legal systems and sovereign risk; increased scrutiny of the health care industry by government agencies; and the Company's ability to successfully separate the Company's Orthopaedics business and realize the anticipated benefits from the planned separation. A further list and descriptions of these risks, uncertainties and other factors can be found in Johnson & Johnson's most recent Annual Report on Form 10-K, including in the sections captioned “Cautionary Note Regarding Forward-Looking Statements” and “Item 1A. Risk Factors,” and in Johnson & Johnson's subsequent Quarterly Reports on Form 10-Q and other filings with the Securities and Exchange Commission. Copies of these filings are available online at www.sec.gov, www.jnj.com, investor.jnj.com, or on request from Johnson & Johnson. Any forward-looking statement made in this release speaks only as of the date of this release. Johnson & Johnson does not undertake to update any forward-looking statement as a result of new information or future events or developments.

Cautionary note on Non-GAAP financial measures

This presentation refers to certain non-GAAP financial measures. These non-GAAP financial measures should not be considered replacements for, and should be read together with, the most comparable GAAP financial measures.

A reconciliation of these non-GAAP financial measures to the most directly comparable GAAP financial measures can be found in the accompanying financial schedules of the earnings release and the Investor Relations section of the Company's website.

Strategic partnerships, collaborations & licensing arrangements

During the course of this presentation, we will discuss a number of products and compounds developed in collaboration with strategic partners or licensed from other companies. The following is an acknowledgement of those relationships:

Oncology

IMBRUVICA developed in collaboration and co-marketed in the U.S. with Pharmacyclics, LLC, an AbbVie company; ZYTIGA licensed from BTG International Ltd.; VELCADE developed in collaboration with Millennium: The Takeda Oncology Company; DARZALEX and DARZALEX FASPRO licensed from Genmab A/S; BALVERSA licensed and discovered in collaboration with Astex Pharmaceuticals, Inc.; ERLEADA licensed from Regents of California and Memorial Sloan Kettering; CARVYKTI licensed and developed in collaboration with Legend Biotech USA Inc. and Legend Biotech Ireland Limited; AKEEGA licensed from TESARO, Inc., an oncology-focused business within GSK, and from BTG International Ltd.; RYBREVANT developed under license with Genmab A/S; LAZCLUZE licensed from Yuhan Corporation; DuoBody platform licensed from Genmab A/S relates to several bispecific antibody programs; OMT animal platform licensed from OMT Inc. relates to several antibody programs; ENHANZE platform licensed from Halozyme Therapeutics, Inc.; bleximenib R&D co-funding agreement with Blackstone Life Sciences

Immunology

REMICADE and SIMPONI/ SIMPONI ARIA marketing partners are Schering-Plough (Ireland) Company, a subsidiary of Merck & Co., Inc. and Mitsubishi Tanabe Pharma Corporation; TREMFYA discovered using MorphoSys AG antibody technology; ICOTYDE was discovered through a collaboration with Protagonist Therapeutics – Janssen retains exclusive rights to develop and commercialize for a broad range of indications; JNJ-4804 R&D co-funding agreement with Royalty Pharma plc

Neuroscience

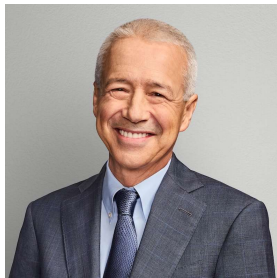
INVEGA SUSTENNA/ XEPLION/ INVEGA TRINZA/ TREVICTA/ INVEGA HAFYERA/ BYANLI are subject to a technology license agreement from Alkermes Pharma Ireland Limited, and RISPERDAL CONSTA developed in collaboration with Alkermes, Inc.

Other

PREZCOBIX / REZOLSTA fixed-dose combination, SYMTUZA and ODEFSEY developed in collaboration with Gilead Sciences, Inc., and JULUCA and CABENUVA developed in collaboration with ViiV Healthcare UK., INVOKANA/ INVOKAMET/ VOKANAMET/ INVOKAMET XR fixed-dose combination licensed from Mitsubishi Tanabe Pharma Corporation; XARELTO co-developed with Bayer HealthCare AG; PROCIT/ EPREX licensed from Amgen Inc.; UPTRAVI license and supply agreement with Nippon Shinyaku (co-promotion in Japan), and OPSUMIT co-promotion agreement with Nippon Shinyaku in Japan

Agenda

- 1 CEO Remarks
- 2 Sales performance and earnings review
- 3 Capital allocation and guidance update
- 4 Q&A



Joaquin Duato
Chairman and
Chief Executive Officer



Joseph J. Wolk
Executive Vice President,
Chief Financial Officer



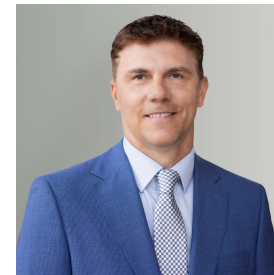
Jennifer Taubert
Executive Vice President,
Worldwide Chairman,
Innovative Medicine



John Reed
Executive Vice President,
Innovative Medicine, R&D



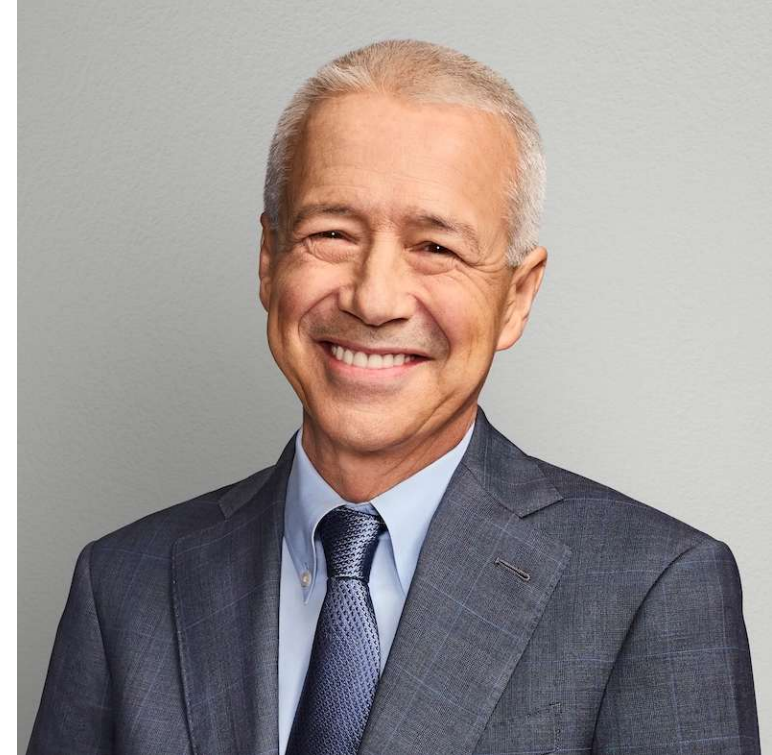
Tim Schmid
Executive Vice President,
Worldwide Chairman,
MedTech



Darren Snellgrove
Vice President,
Investor Relations

Joaquin Duato

Chairman and Chief Executive Officer

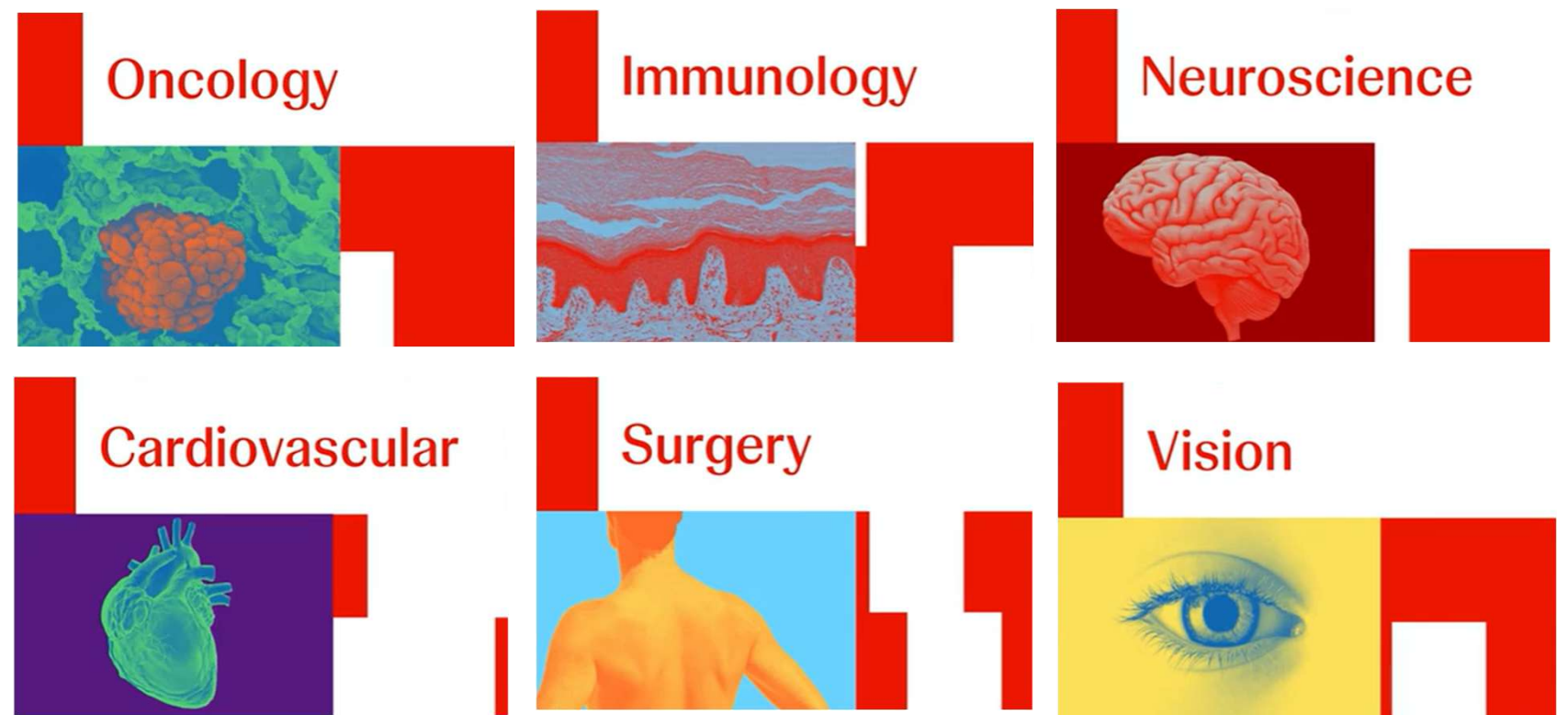


Q1 Earnings Summary

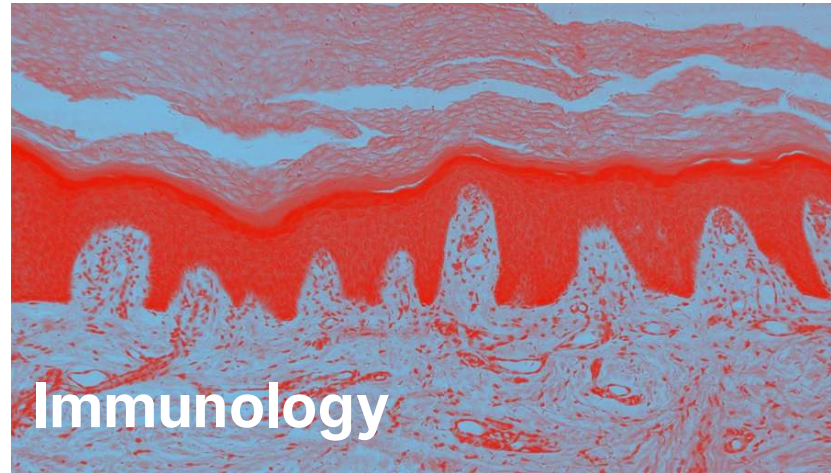
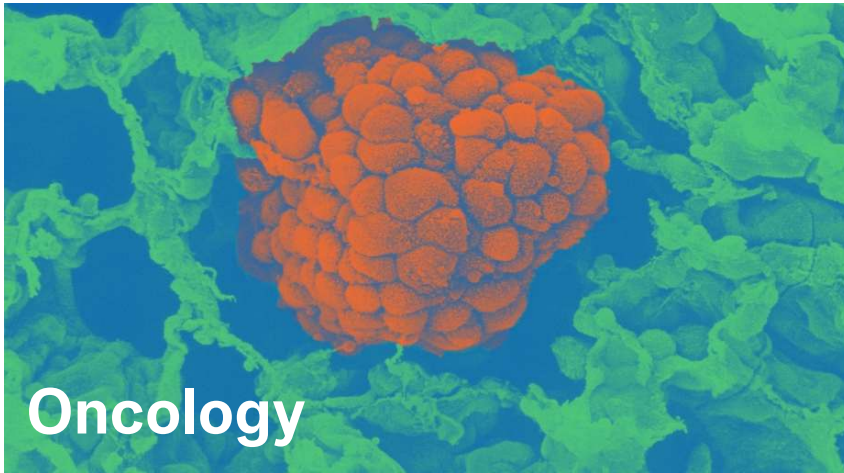
6.4%

operational sales
growth^{1,2}

Six key businesses driving growth



Innovative Medicine: 7.4% operational sales growth^{1,2}



ONCE-DAILY Akeega[®] (niraparib/abiraterone acetate) 100mg/500mg tablets + 50mg/500mg tablets
 Balversa[™] (erdafitinib) tablets
 CARVYKTI[®] (ciltacabtagene autoleucl) for injection
 DARZALEX Faspro[®] (daratumumab and hyaluronidase-fihj) Injections for subcutaneous use: 1.1 g/100mL/500 mg/mL
 Erleada[®] (apalutamide) 60mg tablets / 240mg tablet
 imbruvica[®] (ibrutinib)
 inlexzo[®] gemcitabine intravesical system
 RYBREVANT Faspro[™] + LAZCLUZE[™] (amivantamab and hyaluronidase) (lazertinib) Tablets 100mg/100mg
 TALVEY[™] (talquetamab-tgvs) Injections for subcutaneous use 2 mg/mL and 40 mg/mL
 TECVAYLI[™] (teclistamab)
 Zytiga[®] abiraterone acetate

ICOTYDE[™] (icotrokinra) tablets
 Tremfya[®] (guselkumab)
 imaavy[™] (nipocalimab-aahu)
 Simponi[®] golimumab
 Simponi ARIA[®] golimumab for infusion
 Remicade[®] INFLIXIMAB
 Stelara[®] (ustekinumab)

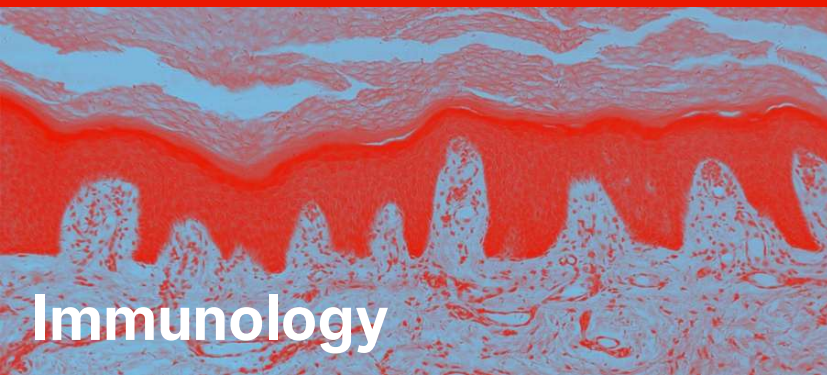
CAPLYTA[®] (lumateperone) capsules 42 mg
 INVEGA SUSTENNA[®] paliperidone palmitate extended-release injectable suspension
 INVEGA TRINZA[®] paliperidone palmitate extended-release injectable suspension 273 mg, 410 mg, 546 mg, 819 mg
 Spravato[™] (esketamine) [®] nasal spray
 INVEGA HAFYERA[™] paliperidone palmitate extended-release injectable suspension 1092mg, 1560mg

¹ Non-GAAP measure; excludes the impact of translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)

² Includes an approximate (920) basis point headwind from STELARA

Note: Includes existing products and planned future launches

ICOTYDE™ U.S. FDA approval brings new option for first-line systemic treatment of plaque psoriasis

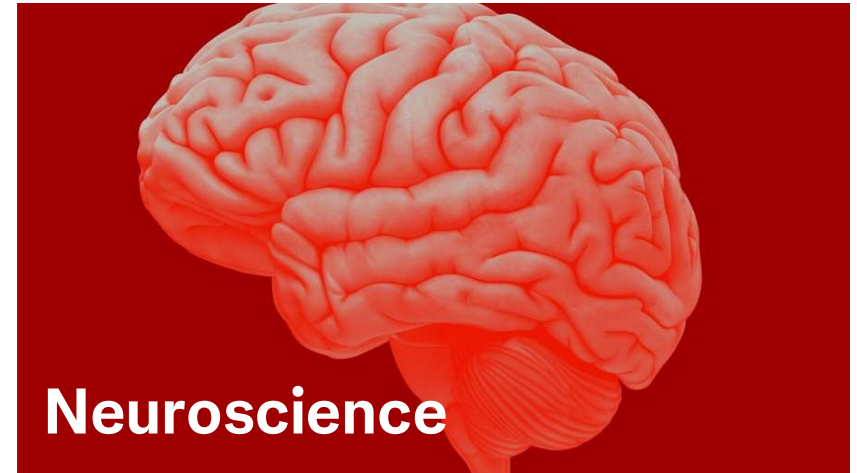
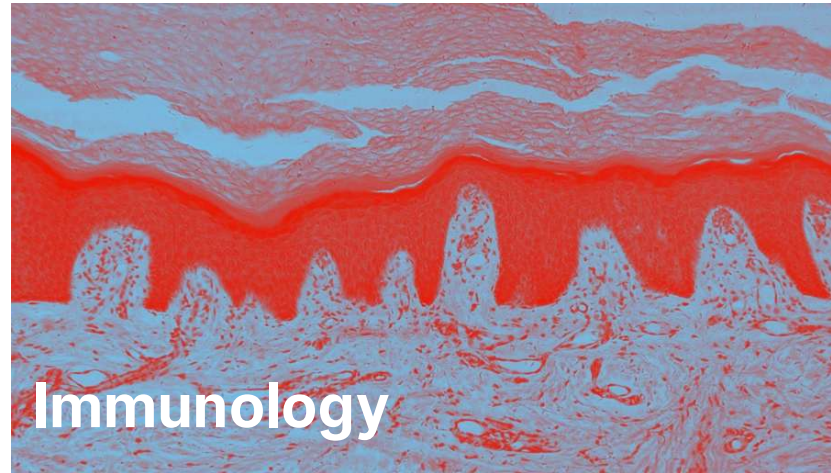
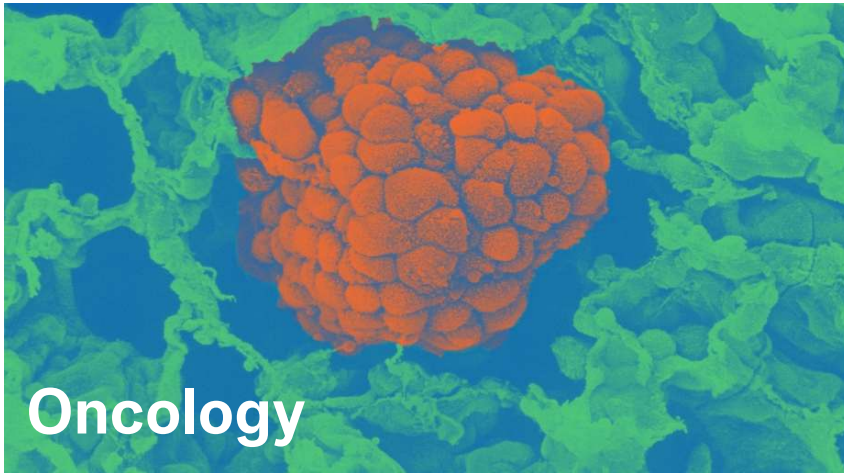


First and only IL-23R targeted oral peptide to deliver complete skin clearance and favorable safety profile in a once-daily pill

- ✓ Preferred first line systemic treatment for patients 12 and older with moderate to severe plaque psoriasis
- ✓ No mandatory TB test or lab monitoring
- ✓ Full launch occurred immediately on approval
- ✓ Potential to revolutionize how psoriatic disease is treated



Innovative Medicine: 7.4% operational sales growth^{1,2}



ONCE-DAILY Akeega® (niraparib/abiraterone acetate) 100mg/500mg tablets + 50mg/500mg tablets
 Balversa™ (erdafitinib) tablets
 CARVYKTI® (ciltacabtagene autoleucel)
 DARZALEX Faspro® (daratumumab and hyaluronidase-fihj) Injections for subcutaneous use, 1.1 g/100mL/500 mg/mL
 Erleada® (apalutamide) 60mg tablets / 240mg tablet
 imbruvica® (ibrutinib)
 inlexzo® gemcitabine intravesical system
 RYBREVANT Faspro™ + LAZCLUZE™ (amivantamab and hyaluronidase) (lazertinib) Tablets 100mg/500mg, 1.24g/500mg
 TALVEY™ (talquetamab-tgvs) Injections for subcutaneous use 2 mg/mL and 40 mg/mL
 TECVAYLI™ (teclistamab)
 Zytiga® abiraterone acetate

ICOTYDE™ (icotrokinra) tablets
 Tremfya® (guselkumab)
 imaavy™ (nipocalimab-aahu)
 Simponi® golimumab
 Simponi ARIA® golimumab for infusion
 Remicade® INFLIXIMAB
 Stelara® (ustekinumab)

CAPLYTA® (lumateperone) capsules 42 mg
 INVEGA SUSTENNA® paliperidone palmitate extended-release injectable suspension
 INVEGA TRINZA® paliperidone palmitate extended-release injectable suspension 273 mg, 410 mg, 546 mg, 819 mg
 Spravato™ (esketamine)® nasal spray
 INVEGA HAFYERA™ paliperidone palmitate extended-release injectable suspension 1092mg, 1560mg

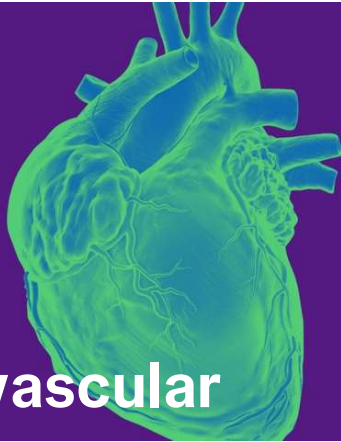
¹ Non-GAAP measure; excludes the impact of translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)

² Includes an approximate (920) basis point headwind from STELARA

Note: Includes existing products and planned future launches

MedTech: 4.6% operational sales growth¹

Cardiovascular



VARIPULSE™ Platform



Dual Energy THERMOCOOL
SMARTTOUCH™ SF Catheter



Shockwave Intravascular
Lithotripsy System

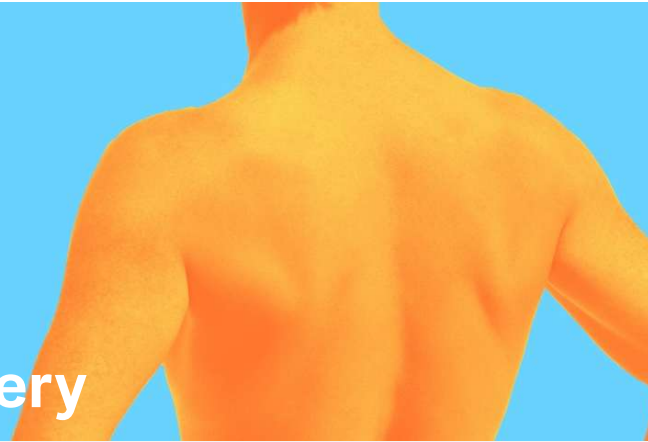


Impella® Heart
Pump Technology



OMNYPULSE™
Catheter

Surgery



ETHICON™ 4000
Surgical Stapler



MONARCH™
Platform



OTTAVA™
Robotic
Surgical
System

Vision



ACUVUE® OASYS 1-Day Family



TECNIS Odyssey™



TECNIS PureSee™

2026 will be a year
of accelerated impact and
growth

Darren Snellgrove

Vice President,
Investor Relations



1st Quarter 2026 sales

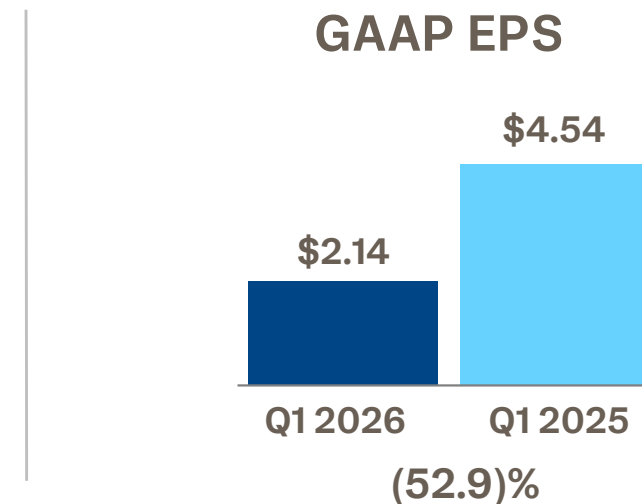
Dollars in billions Regional sales results	Q1 2026	Q1 2025	% Change	
			Reported	Operational ¹
U.S.	\$13.3	\$12.3	8.3%	8.3%
Europe	5.8	5.1	14.5	2.7
Western Hemisphere (ex U.S.)	1.3	1.2	10.8	2.5
Asia-Pacific, Africa	3.6	3.3	8.5	6.1
International	10.7	9.6	11.9	3.9
Worldwide (WW)	\$24.1	\$21.9	9.9%	6.4%

J&J ¹ Non-GAAP measure; excludes the impact of translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)
 Note: Values may be rounded

1st Quarter 2026 financial highlights

Dollars in billions, except EPS

Reported %; **Operational %**¹



J&J ¹Non-GAAP measure; excludes the impact of translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)

²Non-GAAP measure; excludes intangible amortization expense and special items; see reconciliation schedules on the Investor Relations section of the [company's website](#)

Innovative Medicine highlights – 1st quarter 2026

Strong operational growth¹ of 7.4% driven primarily by Oncology and Neuroscience

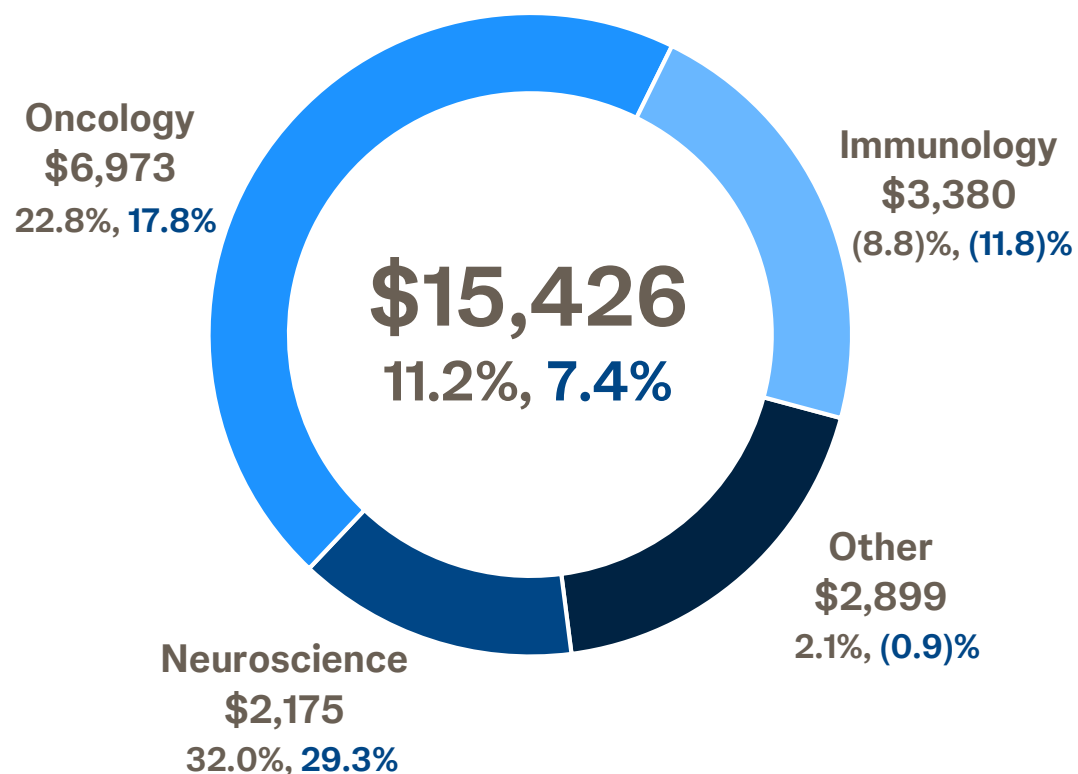
Stelara impacted results¹ by ~(920) basis points

Reported: WW 11.2%, U.S. 9.6%, Int'l 13.4%

Operational¹: WW 7.4%, U.S. 9.6%, Int'l 4.3%

WW sales \$MM

■ Reported growth ■ Operational growth¹



Key drivers of operational performance¹

Oncology	<ul style="list-style-type: none"> DARZALEX increase primarily driven by strong share gains and market growth, partially offset by inventory dynamics CARVYKTI increase driven by share gains and continued site expansion TECVAYLI growth driven by launch uptake and share gains from expansion in the community setting and recent U.S. TECVAYLI + DARZALEX FASPRO approval TALVEY growth driven by share gains from expansion in the community setting RYBREVANT/LAZCLUZE growth driven by launch uptake and share gains ERLEADA increase driven by continued share gains in mCSPC and market growth IMBRUVICA decrease driven by share loss due to continued competitive pressure and unfavorable patient mix
Immunology	<ul style="list-style-type: none"> TREMFYA growth due to share gains across all indications with significant IBD launch momentum and market growth SIMPONI/SIMPONI ARIA and REMICADE decrease driven by share loss, biosimilar competition, and unfavorable patient mix, partially offset by market growth STELARA decline driven by the impact of biosimilar competition, increasing adoption of novel classes, and unfavorable patient mix
Neuroscience	<ul style="list-style-type: none"> SPRAVATO growth driven by continued increased physician and patient demand CAPLYTA driven by strong continued momentum in the aMDD launch INVEGA SUSTENNA / XEPLION / INVEGA TRINZA / TREVICTA increase primarily driven by favorable patient mix
Other	<ul style="list-style-type: none"> UPTRAVI increase driven by market and share growth, partially offset by inventory dynamics OPSUMIT/OPSYNVI growth driven by share gains, market growth, and favorable patient mix PREZISTA / PREZCOBIX / REZOLSTA / SYMTUZA increase driven by favorable patient mix XARELTO decrease driven by continued share erosion

Adjusted operational sales²: WW: 5.6%, U.S. 6.3%, Int'l 4.5%



¹ Non-GAAP measure; excludes the impact of translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)

² Non-GAAP measure; excludes acquisitions and divestitures and translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)

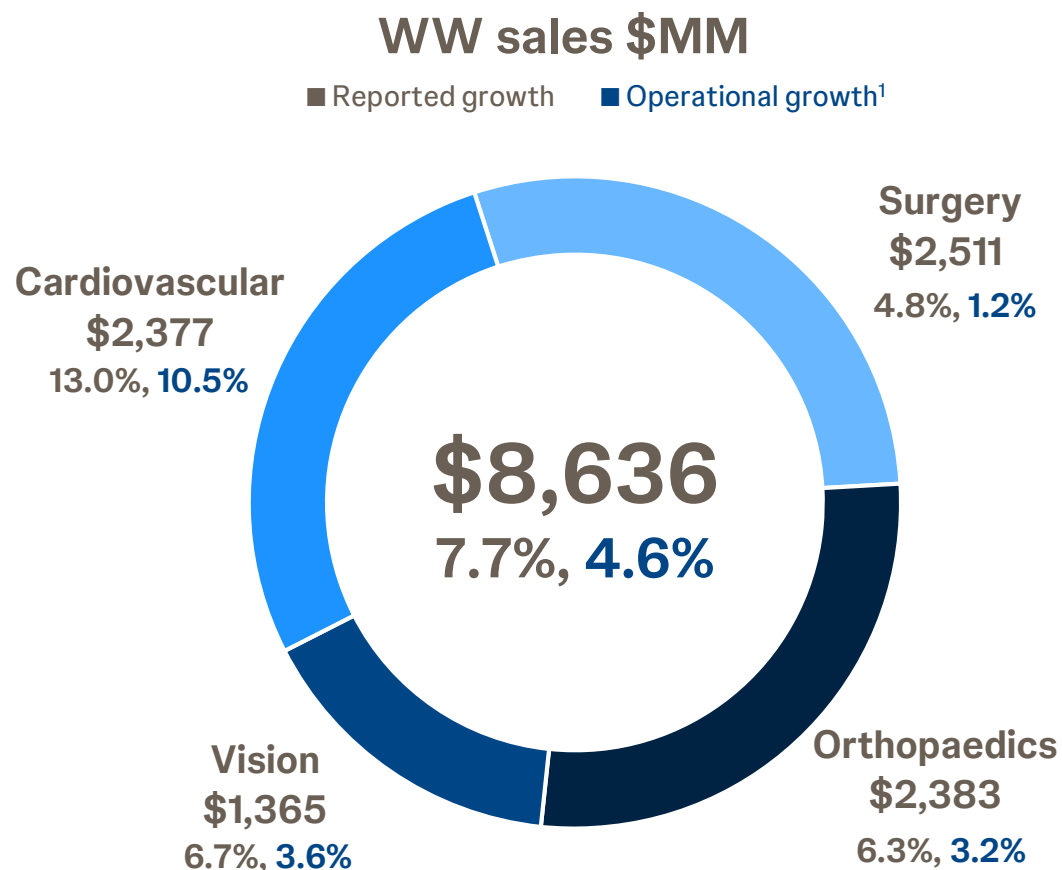
Note: Values may be rounded

MedTech highlights – 1st quarter 2026

Solid operational growth¹ of 4.6% due to commercial execution and innovation

Reported: WW 7.7%, U.S. 5.9%, Int'l 9.7%

Operational¹: WW 4.6%, U.S. 5.9%, Int'l 3.2%



Key drivers of operational performance¹

Cardiovascular	<ul style="list-style-type: none"> • Electrophysiology: Increase driven by procedure growth, commercial execution, new product performance (VARIPULSE, TRUPULSE, NUVISION and CRYSTAL), and OUS inventory dynamics, partially offset by competitive pressures in PFA • Abiomed: Double digit growth driven by continued strong adoption of Impella 5.5 and Impella CP • Shockwave: Double digit growth driven by strong adoption of Coronary and Peripheral portfolios and new product launches (JAVELIN & E8)
Surgery	<ul style="list-style-type: none"> • Advanced: <ul style="list-style-type: none"> • Biosurgery: ~ +5% growth driven by continued strength of the portfolio and commercial execution, partially offset by the impact of the surgery transformation program and VBP in China • Endocutters: ~ -3% due to competitive pressures and VBP in China • Energy: ~ +2% increase driven by new product launches, partially offset by VBP in China • General: Increase primarily due to technology penetration and upgrades within our differentiated Wound Closure portfolio (Barbed & PLUS Sutures) and market expansion, offset by OUS tender timing
Vision	<ul style="list-style-type: none"> • Contact Lenses/Other: Increase driven by strong performance of the ACUVUE OASYS 1-Day family of products and strategic price actions, partially offset by inventory dynamics OUS • Surgical: Growth driven by strength of recent product innovations, robust demand for premium IOLs, and strong commercial execution, partially offset by competitive pressures in the U.S.
Orthopaedics	<p>Growth across all platforms primarily driven by new product launches and strong commercial execution:</p> <ul style="list-style-type: none"> • Hips: Increase driven by new product launches (EMPHASYS) • Knees: Increase driven by strength of the ATTUNE portfolio, in part driven by pull through related to the VELYS Robotic assisted solutions • Trauma: Growth driven by recently launched products (VOLT) • Spine, Sports & Other: New product innovations (TriAltis) and growth in shoulders, offset by competitive pressures and inventory dynamics

Adjusted operational sales²: WW 4.7%, U.S. 6.1%, Int'l 3.2%



¹ Non-GAAP measure; excludes the impact of translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#)

² Non-GAAP measure; excludes acquisitions and divestitures and translational currency; see reconciliation schedules on the Investor Relations section of the [company's website](#).

Note: Values may be rounded

Condensed consolidated statement of earnings

1st Quarter 2026

(Unaudited; Dollar and shares in millions except per share figures)

	2026		2025		% Increase (Decrease)
	Amount	% to Sales	Amount	% to Sales	
Sales to customers	\$24,062	100.0	\$21,893	100.0	9.9
Cost of products sold	8,106	33.7	7,357	33.6	10.2
Gross Profit	15,956	66.3	14,536	66.4	9.8
Selling, marketing and administrative expenses	6,034	25.1	5,112	23.3	18.0
Research and development expense	3,527	14.7	3,225	14.7	9.4
In-process research and development impairments	36	0.1	-	-	
Interest (income) expense, net	43	0.2	(128)	(0.6)	
Other (income) expense, net	294	1.2	(7,321)	(33.4)	
Restructuring	32	0.1	17	0.1	
Earnings before provision for taxes on income	5,990	24.9	13,631	62.3	(56.1)
Provision for taxes on income	755	3.1	2,632	12.1	(71.3)
Net Earnings	\$5,235	21.8	\$10,999	50.2	(52.4)
Net earnings per share (Diluted)	\$2.14		\$4.54		(52.9)
Average shares outstanding (Diluted)	2,445.2		2,423.8		
Effective tax rate	12.6%		19.3%		
Adjusted earnings before provision for taxes and net earnings¹					
Earnings before provision for taxes on income	\$7,821	32.5	\$8,011	36.6	(2.4)
Net earnings	\$6,614	27.5	\$6,706	30.6	(1.4)
Net earnings per share (Diluted)	\$2.70		\$2.77		(2.5)
Effective tax rate	15.4%		16.3%		

J&J ¹ Non-GAAP measure; excludes intangible amortization expense and special items; see reconciliation schedules on the Investor Relations section of the [company's website](#)

Adjusted earnings before provision for taxes on income by segment

1st Quarter 2026

(Unaudited; Dollar in millions)

Innovative Medicine

	2026		2025		% Increase (Decrease)
	Amount	% to Sales	Amount	% to Sales	
Sales to customers	\$15,426	100.0	\$13,873	100.0	11.2
Cost of products sold	3,617	23.4	3,371	24.3	7.3
Gross Profit	\$11,809	76.6	10,502	75.7	12.4
Selling, marketing and administrative expenses	2,918	18.9	2,261	16.3	29.1
Research and development expense	2,813	18.2	2,548	18.4	10.4
Other segment items ¹	(43)	(0.2)	(204)	(1.5)	
Adjusted segment income before tax ²	\$6,121	39.7	5,897	42.5	3.8

MedTech

	2026		2025		% Increase (Decrease)
	Amount	% to Sales	Amount	% to Sales	
Sales to customers	\$8,636	100.0	8,020	100.0	7.7
Cost of products sold	3,192	37.0	2,795	34.8	14.2
Gross Profit	\$5,444	63.0	5,225	65.2	4.2
Selling, marketing and administrative expenses	2,906	33.6	2,656	33.1	9.4
Research and development expense	714	8.3	671	8.4	6.4
Other segment items ¹	(98)	(1.2)	(182)	(2.2)	
Adjusted segment income before tax ²	\$1,922	22.3	2,080	25.9	(7.6)

Enterprise

	2026		2025		% Increase (Decrease)
	Amount	% to Sales	Amount	% to Sales	
Adjusted income before tax ²	\$7,821	32.5	\$8,011	36.6	(2.4)

¹ Includes other Income and Expense

² Non-GAAP measure; excludes intangible amortization expense and special items; see reconciliation schedules on the Investor Relations section of the [company's website](#)

Note: For expenses not allocated to segments, see reconciliation schedules on the Investor Relations section of the [company's website](#)

Joseph J. Wolk

Executive Vice President,
Chief Financial Officer



Capital allocation strategy



Dollars in billions	Q1 2026
Cash and marketable securities	\$22.1
Debt	(\$55.0)
Net debt	(\$32.9)
Free cash flow ^{1,2}	~\$1.5

Note: Values may be rounded

Q1 2026:

\$3.5B invested in R&D

\$3.1B in dividends paid to shareholders

Note: Values may be rounded

J&J ¹ Non-GAAP measure; defined as cash flow from operating activities, less additions to property, plant and equipment
² Estimated as of April 14, 2026. Cash flow from operations, the most directly comparable GAAP financial measure, will be included in subsequent SEC filings

2026 P&L guidance

Operational² sales guidance of 6.4% and adjusted operational EPS^{2,4} at 5.7% (midpoints)

	April 2026	January 2026	Comments
Adjusted operational sales ^{1,2}	5.6% - 6.6%	5.4% - 6.4%	Increasing midpoint to 6.1%
Operational sales ²	\$99.7B - \$100.7B 5.9% - 6.9%	\$99.5B - \$100.5B 5.7% - 6.7%	Increasing midpoint by \$0.2B to 6.4%
Estimated reported sales ³	\$100.3B - \$101.3B 6.5% - 7.5%	\$100.0B - \$101.0B 6.2% - 7.2%	Increasing midpoint by \$0.3B to 7.0% Incremental FX impact of \$0.1B or 0.1%
Adjusted pre-tax operating margin ^{4,5}	Increase of at least 50 bps	Increase of at least 50 bps	Maintaining
Net other income ⁴	\$1.0 - \$1.2 billion	\$1.0 - \$1.2 billion	Maintaining
Net interest expense / (income)	\$300 - \$400 million	\$300 - \$400 million	Maintaining
Effective tax rate ⁴	17.5% - 18.5%	17.5% - 18.5%	Maintaining
Adjusted EPS (operational) ^{2,4}	\$11.30 - \$11.50 4.7% - 6.7%	\$11.28 - \$11.48 4.5% - 6.5%	Increasing midpoint by \$0.02 to 5.7%
Adjusted EPS (reported) ^{3,4}	\$11.45 - \$11.65 6.1% - 8.1%	\$11.43 - \$11.63 5.9% - 7.9%	Increasing midpoint by \$0.02 to 7.1% Maintaining FX impact of \$0.15 or 1.4%

¹ Non-GAAP measure; excludes acquisitions and divestitures

² Non-GAAP measure; excludes the impact of translational currency

³ Calculated using Euro Average Rate: April 2026 = \$1.17 and January 2026 = \$ 1.17

Note: Values may be rounded

⁴ Non-GAAP measure; excludes intangible amortization expense and special items

⁵ Sales less: COGS, SM&A and R&D expenses

Phasing Considerations

Anticipate fairly consistent operational¹ sales growth in the first and second half; 53rd week impacts the second half

Innovative Medicine

- Expect more pronounced impact from newly launched products as the year progresses
- Anticipate generic competition for OPSUMIT (second half in U.S.) and SIMPONI (first half in EU; potentially second half in U.S.)
- Expect impact of voluntary agreement with the U.S. government to be evenly distributed throughout the year

MedTech

- Expect continued acceleration of newly launched products as the year progresses with normalized seasonality
- Surgery Transformation sales headwinds expected to increase through the year
- Anticipate additional rounds of VBP in China throughout the year, heavier in the second half of the year

P&L

- Expect heavier investment in the first half of the year
- Intra-Cellular benefit laps in Q2
- Anticipate higher earnings per share growth in the second half of the year compared to the first half

Anticipated 2026 milestones¹ driving long-term value creation

Innovative Medicine

ICOTYDE in PsO

TECVAYLI + DARZALEX in RRMM

TREMFYA in PsA SD

TECVAYLI in RRMM

INLEXZO in HR NMIBC

nipocalimab in WAIHA and SLE

CAPLYTA in bipolar mania

JNJ-4804

ERLEADA in LPC & HRPC

MedTech

OTTAVA

ETHIZIA

STSF Dual Energy Catheter

C2 Aero Catheters

TECNIS PureSee IOL

ATTUNE Hinge

VOLT

MONARCH Urology

VARIPULSE Pro

Save the Date

Johnson & Johnson

Enterprise Business Review

Tuesday, December 8, 2026

**Innovative
Medicine**

Oncology



Immunology



Neuroscience



MedTech

Surgery



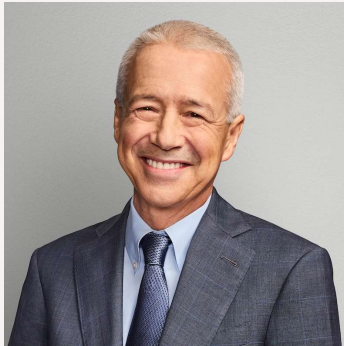
Cardiovascular



Vision



Q&A



Joaquin Duato
Chairman and
Chief Executive Officer



Jennifer Taubert
Executive Vice President,
Worldwide Chairman,
Innovative Medicine



Tim Schmid
Executive Vice President,
Worldwide Chairman,
MedTech



Joseph J. Wolk
Executive Vice President,
Chief Financial Officer



John Reed
Executive Vice President,
Innovative Medicine, R&D



Darren Snellgrove
Vice President,
Investor Relations

Johnson & Johnson

Johnson & Johnson Innovative Medicine Pipeline

Key Events in 2026*

POTENTIAL APPROVALS US/EU

US	TREMFYA (guselkumab) Psoriatic Arthritis Structural Damage (APEX)
✓ US EU	ICOTYDE (icotrokinra) Psoriasis (ICONIC)
EU	AKEEGA (niraparib/abiraterone) M1 Metastatic Castration-Sensitive Prostate Cancer (AMPLITUDE)
✓ US	DARZALEX (daratumumab) Frontline multiple myeloma transplant ineligible (CEPHEUS)
✓ US EU	TECVAYLI (teclistamab) Multiple Myeloma 1-3PLs (MajesTEC-3)
EU	TECVAYLI (teclistamab) Relapsed Refractory Multiple Myeloma CD38 exposed (MajesTEC-9)

PLANNED SUBMISSIONS US/EU

✓ US	nipocalimab Warm Autoimmune Hemolytic Anemia (ENERGY)
US	CAPLYTA (lumateperone) Bipolar Mania
US EU	ERLEADA (apalutamide) Localized Prostate Cancer (ATLAS)
US EU	ERLEADA (apalutamide) High Risk Prostate Cancer (PROTEUS)
US	bleximenib Relapsed Refractory Acute Myeloid Leukemia (cAMeLot-1)
✓ US ✓ EU	TALVEY (talquetamab) Relapsed Refractory Multiple Myeloma A-CD38 Naïve (MonumenTAL-3)
✓ US ✓ EU	TECVAYLI (teclistamab) Relapsed Refractory Multiple Myeloma CD38 exposed (MajesTEC-9)
✓ EU	TECVAYLI (teclistamab) Multiple Myeloma 1-3PLs (MajesTEC-3)
US EU	INLEXZO (gemcitabine intravesical delivery system) High Risk Non Muscle Invasive Bladder Cancer BCG Experienced (SunRISe-5)

POTENTIAL CLINICAL DATA PRESENTATIONS¹

Phase III

✓	ICOTYDE (icotrokinra) Psoriasis (ICONIC-ADVANCE 1&2 Update)
	ICOTYDE (icotrokinra) Psoriasis (ICONIC-ASCEND)
	icotrokinra Psoriatic Arthritis (ICONIC-PsA)
	ICOTYDE (icotrokinra) Psoriasis (ICONIC-TOTAL Update)
	CAPLYTA (lumateperone) Bipolar Mania (ITI-007-452)
	nipocalimab Warm Autoimmune Hemolytic Anemia (ENERGY)

Phase I/ II

	ERLEADA (apalutamide) Localized Prostate Cancer (ATLAS)	JNJ-4804 Co-antibody Therapy Ulcerative Colitis (DUET-UC)
	ERLEADA (apalutamide) High Risk Prostate Cancer (PROTEUS)	JNJ-4804 Co-antibody Therapy Crohn's Disease (DUET-CD)
	TALVEY (talquetamab) Relapsed Refractory Multiple Myeloma A-CD38 Naïve (MonumenTAL-3)	JNJ-4804 Co-antibody Therapy Psoriatic Arthritis (AFFINITY)
	TECVAYLI (teclistamab) Relapsed Refractory Multiple Myeloma CD38 exposed (MajesTEC-9)	nipocalimab Systemic Lupus Erythematosus (JASMINE)
	INLEXZO (gemcitabine intravesical delivery system) High Risk Non Muscle Invasive Bladder Cancer BCG Experienced (SunRISe-5)	bleximenib Relapsed Refractory Acute Myeloid Leukemia (cAMeLot-1)
		JNJ-1887 sCD59 Geographic Atrophy (PARASOL)

1. In order to be on key events clinical presentation, data must be presented at a major medical meeting.

✓ = Achieved

*This is not a fully exhaustive list of all pipeline programs and assets. The pipeline includes assets currently progressing through clinical trials as well as those under review by regulatory bodies. Inclusion in the pipeline is based on the current status of these programs and assets and does not guarantee continued investment. This information is as of April 14, 2026 to the best of the Company's knowledge. Johnson & Johnson assumes no obligation to update this information.

