

Q2 2026 Results

July 30, 2026

Forward Looking Statements and Non-GAAP Financial Information

This presentation (as well as the oral statements made with respect to information contained herein) contains statements about Bristol-Myers Squibb Company's (the "Company") future financial results, plans, business development strategy, anticipated clinical trials, results and regulatory approvals that constitute forward-looking statements for purposes of the safe harbor provisions under the Private Securities Litigation Reform Act of 1995. All statements that are not statements of historical facts are, or may be deemed to be, forward-looking statements. Actual results may differ materially from those expressed in, or implied by, these statements as a result of various factors, including, but not limited to: (i) new laws, government actions, agreements and regulations, including with respect to pricing controls and market access and the imposition of new tariffs, trade restrictions and export regulations, including the potential for international reference pricing and most-favored nation drug pricing for our products, (ii) our ability to obtain, protect and maintain market exclusivity rights and enforce patents and other intellectual property rights, (iii) our ability to achieve expected clinical, regulatory and contractual milestones on expected timelines or at all, (iv) difficulties or delays in the development and commercialization of new products, (v) difficulties or delays in our clinical trials and the manufacturing, distribution and sale of our products, (vi) adverse outcomes in legal or regulatory proceedings, (vii) risks relating to acquisitions, divestitures, alliances, joint ventures and other portfolio actions and (viii) political and financial instability, including changes in general economic conditions, and (viii) risks related to the use of artificial intelligence or other emerging technologies in various facets of the company's operations, including partnerships related to the use of, or the sharing of such technologies with third parties, which may exacerbate competitive, regulatory, litigation, cybersecurity, and other risk. These and other important factors are discussed in the Company's most recent annual report on Form 10-K and reports on Forms 10-Q and 8-K. These documents are available on the U.S. Securities and Exchange Commission's website, on the Company's website or from Bristol-Myers Squibb Investor Relations. No forward-looking statements can be guaranteed.

In addition, any forward-looking statements and clinical data included herein are presented only as of the date hereof. Except as otherwise required by applicable law, the Company undertakes no obligation to publicly update any of the provided information, whether as a result of new information, future events, changed circumstances or otherwise.

This presentation includes certain non-generally accepted accounting principles ("GAAP") financial measures that we use to describe the Company's performance. The non-GAAP

financial measures are provided as supplemental information and are presented because management has evaluated the Company's financial results both including and excluding the adjusted items or the effects of foreign currency translation, as applicable, and believes that the non-GAAP financial measures presented portray the results of the Company's baseline performance, supplement or enhance management's, analysts' and investors' overall understanding of the Company's underlying financial performance and trends and facilitate comparisons among current, past and future periods. This presentation also provides certain revenues and expenses excluding the impact of foreign exchange ("Ex-FX"). We calculate foreign exchange impacts by converting our current-period local currency financial results using the prior period average currency rates and comparing these adjusted amounts to our current-period results. Ex-FX financial measures are not accounted for according to GAAP because they remove the effects of currency movements from GAAP results.

The non-GAAP information presented herein provides investors with additional useful information but should not be considered in isolation or as substitutes for the related GAAP measures. Moreover, other companies may define non-GAAP measures differently, which limits the usefulness of these measures for comparisons with such other companies. We encourage investors to review our financial statements and publicly filed reports in their entirety and not to rely on any single financial measure. An explanation of these non-GAAP financial measures and a reconciliation to the most directly comparable financial measure are available on our website at www.bms.com/investors.

Also note that a reconciliation of forward-looking non-GAAP measures, including non-GAAP earnings per share (EPS), to the most directly comparable GAAP measures is not provided because comparable GAAP measures for such measures are not reasonably accessible or reliable due to the inherent difficulty in forecasting and quantifying measures that would be necessary for such reconciliation. Namely, we are not, without unreasonable effort, able to reliably predict the impact of accelerated depreciation and impairment charges, legal and other settlements, gains and losses from equity investments and other adjustments. In addition, the Company believes such a reconciliation would imply a degree of precision and certainty that could be confusing to investors. These items are uncertain, depend on various factors and may have a material impact on our future GAAP results.

Certain information presented in the accompanying presentation may not add due to the use of rounded numbers.

Q2 2026 Results



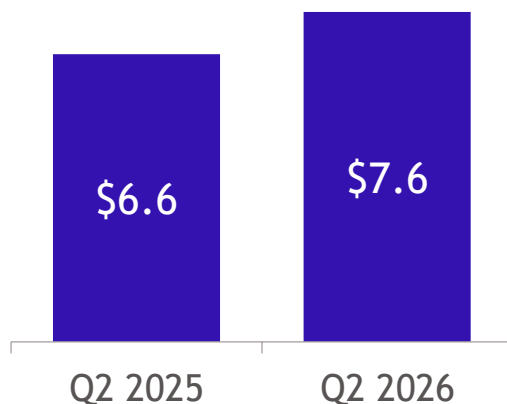
Chris Boerner, PhD
Board Chair and
Chief Executive Officer

Q2 2026 Performance

Growth Portfolio Revenues

\$ in billions

+15%; +14% Ex-FX*



Diverse portfolio of growth drivers

Reblozyl[™]
(luspatercept-aamt)
for injection 25mg + 75mg

Breyanzi[™]
(lisocabtagene maraleucel) SUSPENSION
FOR IV INFUSION

CAMZYOS[™]
(mavacamten) capsules

Opdualag[™]
(nivolumab and relatlimab-mbw)
Injection for intravenous use | 480 mg/160 mg

OPDIVO Qvantig[™]
nivolumab + hyaluronidase-nvhy
SUBCUTANEOUS INJECTION | 120 mg + 2,000 units / mL

Key Milestones¹

- **ASCO**
Data highlighted the breadth of innovation across our oncology franchise
- **CELMoDs: advancing towards commercialization**
Iberdomide PDUFA: August 17, 2026
Mezigdomide PDUFA: May 13, 2027
- **Iza-bren**
Initiating 4th global pivotal trial in 1L EGFRmut NSCLC
- **Pumitamig**
Initiating Phase 2 novel combination study with imzokitug

*See "Forward-Looking Statements and Non-GAAP Financial Information" 1. Not an exhaustive list of assets, programs, or indications

Significant data expected in 2026...and beyond*

NME registrational data

2026

- Admilparant **IPF** (ALOFT-IPF)
- Arlo-cel **4L+ MM** (QUINTESSENTIAL)
- Iberdomide **RRMM** PFS (EXCALIBER-RRMM)
- Mezigdomide **RRMM** (SUCCESSOR-2) (Mar'26)
- Milvexian **SSP** (LIBREXIA-STROKE¹)
- RYZ101 **2L+ GEP-NETs** (ACTION-1)

2027

- Golcadomide **High-Risk 1L LBCL** (GOLSEEK-1)
- Milvexian **AF** (LIBREXIA-AF¹)
- Zola-cel **SLE** (Breakfree-SLE)

2028

- Atigotatug + nivolumab **1L ES-SCLC** (TIGOS)
- Iza-bren **1L TNBC** (IZABRIGHT-Breast01)
- Pumitamig **1L ES-SCLC** (ROSETTA-Lung-01²)

LCM pivotal data

2026

- Sotyktu **SLE** (POETYK SLE-1 & 2)

2027

- Admilparant **PPF** (ALOFT-PPF)
- Cobenfy **AD Psychosis** (ADEPT)
- Cobenfy **Bipolar-I** (BALSAM-1 & 2)
- Mezigdomide **RRMM** (SUCCESSOR-1)
- Sotyktu **Sjogren's Disease** (POETYK SjS-1)

2028

- Arlo-cel **2-4L MM** (QUINTESSENTIAL-2)
- Cobenfy **AD Agitation** (ADAGIO-2)
- Cobenfy **AD Cognition** (MINDSET-1 & 2)
- Cobenfy **Adjunctive Bipolar-I** (BALSAM-4)
- Golcadomide **2L+ FL** (GOLSEEK-4)
- Iza-bren **EGFRm NSCLC** (IZABRIGHT-Lung01)
- Krazati **1L NSCLC** (KRYSTAL-4)
- Krazati **1L NSCLC PD-L1 ≥50%** (KRYSTAL-7)
- Reblozyl **1L NTD MDS Associated Anemia** (ELEMENT)
- Zola-cel **SSc** (Breakfree-SSc)

Key next wave early-stage data

2026

- BCMaXGPRC5D dual-targeting CAR T **RRMM**
- Golcadomide **1L FL** (GOLSEEK-2)
- Imzokitug (anti-CCR8) **Solid Tumors**
- Delocamten (MYK-224) **HFpEF** (AURORA)
- Navlimetostat (PRMT5 inhibitor) **Solid Tumors**
- Pumitamig **Solid Tumors²**
- Zola-cel **Autoimmune Diseases** (Breakfree-1 & 2)

2027

- Moponetug (anti-MTBR-tau) **Alzheimer's Disease** (TargetTau-1)
- Irafamdastat (FAAH/MAGL) **AD Agitation** (BALANCE-AAD-1)
- Irafamdastat (FAAH/MAGL) **MS Spasticity** (BALANCE-MSS-1)

*See "Forward-Looking Statements and Non-GAAP Financial Information", NME: New Molecular Entity, LCM: Life Cycle Management; 1. Trial conducted by Johnson & Johnson; 2. Trial conducted by BioNTech. Studies shown with dates in parenthesis have reported readouts

Rewiring how we operate*



AI-Powered Execution

Expanding AI to help teams move faster and operate more efficiently, further strengthened through our Anthropic & NVIDIA partnerships



Operating Discipline

Company-wide savings through strategic productivity initiative, combined with strong free cash flow and balance sheet strength



Financial Flexibility

Invest in growth drivers, pursue strategic business development opportunities, and return cash to shareholders

*See “Forward-Looking Statements and Non-GAAP Financial Information”

Q2 2026 Results

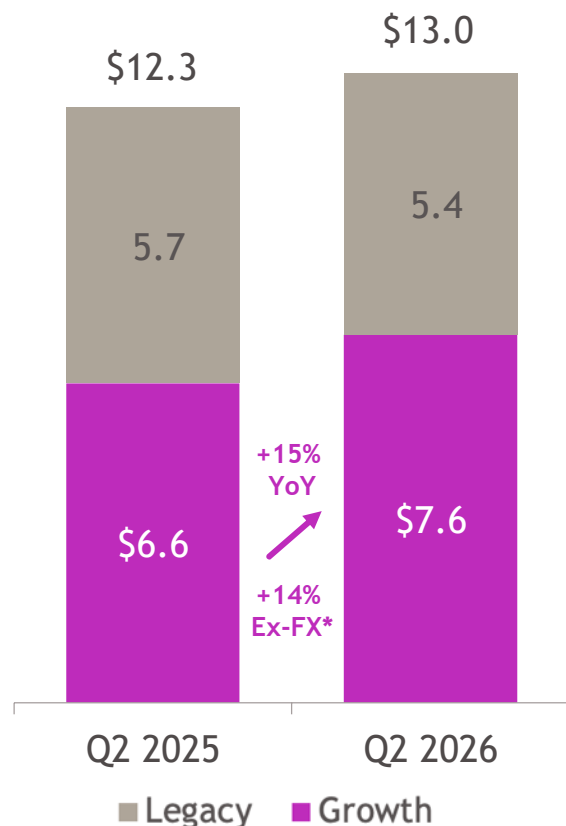


David Elkins

Executive Vice President
and Chief Financial Officer

Growth Portfolio continues to drive revenues

\$ in billions



Growth Portfolio

OPDIVO
(nivolumab)
INJECTION FOR INTRAVENOUS USE 10 mg/mL

OPDIVO Qvantig
nivolumab + hyaluronidase-nvhy
SUBCUTANEOUS INJECTION 100 mg + 2,000 units / mL

ORENCIA
(abatacept)
INJECTION FOR INTRAVENOUS USE 100 mg/mL

SOTYKTU
(deucravacitinib) 6 mg tablets

Opdualag
(nivolumab and relatlimab-mbaw)
INJECTION FOR INTRAVENOUS USE 480 mg/160 mg

YERVOY
(ipilimumab)
INJECTION FOR INTRAVENOUS INFUSION

ZEPOSIA
(ozanimod) 0.2 mg capsules

KRAZATI
(adagrasib) 200 mg tablets

COBENFY
(xonometrine and tropism chloride) capsules
50 mg/20 mg, 100 mg/20 mg, 125 mg/30 mg

CAMZYOS
(mavacamten) capsules

Breyanzi
(lisocabtagene maraleucel) SUSPENSION FOR INTRAVENOUS INFUSION

Reblozyl
(luspatercept-aamt) for injection 25 mg + 75 mg

Other Growth Brands¹

Legacy Portfolio

Eliquis
(apixaban) tablets 5 mg, 2.5 mg

Revlimid
(lenalidomide) capsules
2.5, 5, 10, 15, 20, 25 mg

Pomalyst
(pomalidomide) capsules
1, 2, 3, 4 mg

Abraxane
(nanoparticle albumin-bound paclitaxel)

SPRYCEL
dasatinib 100 mg tablets

Other Mature Brands

*See "Forward-Looking Statements and Non-GAAP Financial Information"; 1. Other Growth Brands: Abecma, Augtyro, Emlipicit, Inrebic, Nulojix, Onureg, & Royalty Revenues, including royalties received from Merck on Winrevair

Q2 2026 Oncology product summary

Global Net Sales¹

	\$M	YoY %	Ex-FX* %
 INJECTION FOR INTRAVENOUS USE 10 mg/mL	\$2,485	(3%)	(4%)
 INJECTION FOR INTRAVENOUS INFUSION	\$769	+6%	+5%
 INJECTION FOR INTRAVENOUS USE 480 mg/160 mg	\$349	+23%	+22%
 SUBCUTANEOUS INJECTION 100 mg + 2,000 units/mL	\$261	>200%	>200%
 200 mg TABLETS	\$55	+14%	+14%

Opdivo

- Remains an important revenue contributor to the Growth Portfolio and continues to deliver value to patients

Qvantig

- Continued global growth is driven by meaningful convenience to patients and healthcare systems

Opdualag

- Growth reflects strong demand and established position as a standard of care in 1L melanoma

See "Forward-Looking Statements and Non-GAAP Financial Information"; 1. Abraxane: Q2 2026 WW Sales \$55M - YoY% (47%), (47%) Ex-FX; 2. Opdivo Q2'26 global sales reflect ~\$200M sequential inventory build

Q2 2026 Hematology product summary

Global Net Sales¹

	\$M	YoY %	Ex-FX* %
 (luspatercept-aamt) for injection 25mg + 75mg	\$735	+29%	+29%
 (lisocabtagene maraleucel) SUSPENSION FOR IV INFUSION	\$484	+41%	+41%
 (lenalidomide) capsules	\$425	(49%)	(49%)
 (pomalidomide) capsules	\$204	(71%)	(71%)

Reblozyl

- Growth driven by continued global demand across MDS-associated anemia, with the predominant share in 1L

Breyanzi

- Continued strong demand as the leading CD19 directed CAR T across B-cell malignancies, focused on growing adoption in the community outpatient setting

See “Forward-Looking Statements and Non-GAAP Financial Information”; 1. Abecma: Q2 2026 WW Sales \$72M - YoY% (17%), (16%) Ex-FX; Sprycel: Q2 2026 WW Sales \$88M - YoY% (27%), (26%) Ex-FX*; 2. In the U.S., generic lenalidomide products are no longer volume-limited as of January 31, 2026; 3. In the U.S., generic pomalidomide entered in March of 2026

Q2 2026 Cardiovascular & Immunology product summary

Global Net Sales (Cardiovascular)

	\$M	YoY %	Ex-FX* %
 apixaban	\$4,481	+22%	+21%
 (mavacamten) capsules	\$416	+60%	+59%

Global Net Sales (Immunology)

	\$M	YoY %	Ex-FX* %
 (abatacept)	\$1,034	+7%	+7%
 (deucravacitinib) 6 mg tablets	\$87	+23%	+23%

Camzyos

- Continued strong global demand and market penetration in oHCM
- Strong launch momentum outside the U.S.

Eliquis

- Continued U.S. demand growth and market share gains

Sotyktu

- Continued engagement with the rheumatology community ahead of pivotal SLE and SjD readouts

*See "Forward-Looking Statements and Non-GAAP Financial Information"

Q2 2026 Neuroscience product summary

Global Net Sales

	\$M	YoY %	Ex-FX* %
 ZEPOSIA (ozanimod) 0.22 mg capsules	\$169	+12%	+12%
 COBENFY (xanomeline and trospium chloride) capsules 60mg/20mg, 100mg/20mg, 125mg/30mg	\$63	+81%	+81%

Cobenfy

- Steady growth trajectory expected near term
- Improving patient experience through prescriber education emphasizing optimized dose utilization

*See “Forward-Looking Statements and Non-GAAP Financial Information”

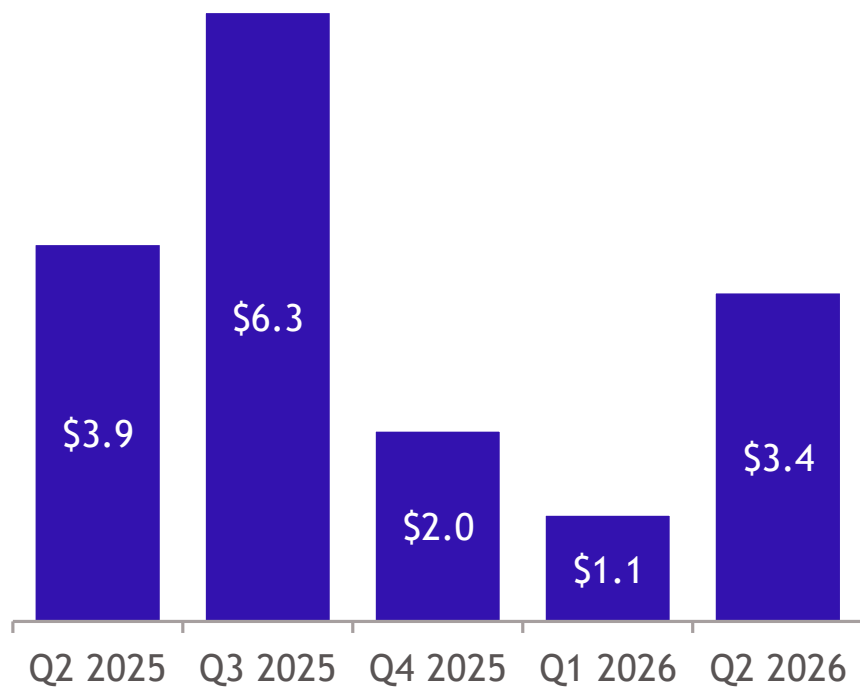
Q2 2026 Financial Performance

\$ in billions, except EPS	GAAP		Non-GAAP*	
	Q2 2026	Q2 2025	Q2 2026	Q2 2025
Total Revenues, net	13.0	12.3	13.0	12.3
Gross Margin % ¹	71.3%	72.5%	71.4%	72.6%
Operating Expenses ²	4.8	4.3	4.1	4.0
Acquired IPR&D	-	1.5	-	1.5
Amortization of Acquired Intangibles	0.4	0.8	-	-
Effective Tax Rate	18.8%	25.9%	16.5%	16.1%
Diluted EPS	1.62	0.64	2.04	1.46
Diluted Shares Outstanding (# in millions)	2,048	2,038	2,048	2,038
Diluted EPS Impact from Acquired IPR&D ³	0.01	(0.57)	0.01	(0.57)

*See “Forward-Looking Statements and Non-GAAP Financial Information”; 1. Gross Margin = Revenue less COGS as a percentage of Revenue; 2. Operating Expenses = SG&A and R&D; 3. Represents the net impact from Acquired IPRD & licensing income

Strategic approach to Capital Allocation

Cash flow from Operations \$B



\$B

Total Cash¹

Total Debt

Q2 2026

~\$11.5

~\$43.1

Business Development

- Pursue opportunities and partnerships to diversify portfolio & strengthen long-term outlook

Balance Sheet Strength

- Strong balance sheet affords financial flexibility
- Maintain strong investment-grade credit rating

Returning Cash to Shareholders

- Remain committed to our dividend²
- ~\$5B share repurchase authorization remaining as of June 30, 2026

1. Cash includes cash, cash equivalents and marketable debt securities; 2. Subject to Board approval

2026 Guidance*

	Non-GAAP ¹	
	April (Prior)	July (Updated)
Total FY Revenues (Reported & Ex-FX)	~\$46.0 - \$47.5B	~\$49.0 - \$50.0B
Gross Margin %	~69-70%	No change
Operating Expenses ²	~\$16.3B	~\$16.5B
Other Income/ (Expense)	~(\$700M)	No change
Tax Rate	~18%	No change
Diluted EPS	\$6.05 - \$6.35	\$6.75 - \$7.00

Diluted weighted-average shares outstanding of 2,050 million were used to calculate 2026 diluted EPS guidance

*The Company does not reconcile forward-looking non-GAAP measures. See “Forward-Looking Statements and Non-GAAP Financial Information”; 2026 Guidance excludes the impact of any potential strategic acquisitions, divestitures, specified items that have not yet been identified and quantified, and the impact of future Acquired IPRD charges and licensing income incurred after June 30, 2026; 1. Guidance provided in April was calculated based on mid-April foreign exchange rates; Guidance provided in July was calculated based on mid-July foreign exchange rates; 2. Operating Expenses = SG&A and R&D

Key Highlights

- FY revenue vs. prior guidance reflects strong first half results and our current projections for the year:
 - Continued Growth Portfolio strength
 - 4% to 6% decline in Legacy Portfolio
 - 20% to 25% growth in WW Eliquis revenue
- Gross margin reflects impact of product mix
- Operating expenses increase reflects growth-oriented initiatives

Q2 2026 Results Q&A



Chris Boerner, PhD
Board Chair,
Chief Executive Officer



David Elkins
Executive VP,
Chief Financial Officer



Adam Lenkowsky
Executive VP,
Chief Commercialization
Officer



Cristian Massacesi, MD
Executive VP,
Chief Medical Officer,
Global Drug Development