

#FutureFresenius: **Biopharma 'Meet the Management'**

Conference call and webcast for investors and analysts

Bad Homburg, 15 December 2025

Safe Harbor Statement

This presentation contains forward-looking statements that are subject to various risks and uncertainties. Future results could differ materially from those described in these forward-looking statements due to a variety of factors, e.g., changes in business, economic, and competitive conditions, regulatory reforms, results of clinical trials, foreign-exchange-rate fluctuations, uncertainties in litigation or investigative proceedings, the availability of financing, and unforeseen impacts of international conflicts.

Financial figures on profit and profitability throughout this presentation, especially EBIT, EBITDA, and related margins, are generally reported "before special items". Hence, these figures exclude certain one-time effects. Regarding the definition of financial performance indicators, these refer to the most recent financial publications available on the Fresenius corporate website.

Fresenius does not undertake any responsibility to update the forward-looking statements contained in this presentation.

Meeting agenda

#	Agenda	Presenter		
I	#FutureFresenius: Rejuvenate in Biopharma	 Michael Sen		
II	Biopharma: Value creation strategy	 Pierluigi Antonelli	 Sang-Jin Pak	
III	Portfolio Rejuvenation for long-term growth	 Fabrice Romanet	 Michael Hammer	
IV	Cost Leadership: Reaping benefits of vertical integration	 Yannick Sorlet	 Jurgen Van Broeck	
V	Commercial Excellence: Balanced footprint across geographies	 Sang-Jin Pak	 Molly Benson	
VI	Q&A	 Michael Sen	 Pierluigi Antonelli	 Sang-Jin Pak
VII	Wrap-up	 Michael Sen		



**#FutureFresenius:
Rejuvenate in Biopharma**

WE ARE COMMITTED TO LIFE

A photograph of a man with short brown hair and a light beard, smiling warmly at the camera. He is sitting on a dark grey couch, holding a sleeping baby in his arms. The baby is wearing a white long-sleeved shirt with thin blue stripes and blue pants. The background is a bright, out-of-focus indoor space with large windows showing greenery outside.

OUR MISSION

We save and improve human lives with affordable, accessible and innovative healthcare products and highest quality in clinical care

OUR VISION

We are the trusted, market-leading healthcare company that unites cutting-edge technology and human care to shape next-level therapies

Addressing three structural healthcare challenges

LONGEVITY GAP

10+ disease-burdened life years

BY 2030, 1.4B PEOPLE WILL BE OVER 60; 84% OF 67M DEATHS EXPECTED TO BE FROM CHRONIC DISEASE¹

“**People are living longer, but spending more years in poor health**

WORKFORCE GAP

>10m health-care worker shortfall

THE WORLD HEALTH ORGANIZATION ESTIMATES A PROJECTED SHORTFALL OF 10 MILLION HEALTH WORKERS BY 2030²

“**There are too few health workers to meet a growing demand for care**

EFFICIENCY GAP

>10% GDP on health expenditure

HEALTH EXPENDITURE IS EXPECTED TO RISE TO MORE THAN 10% OF GLOBAL GDP BY 2030³

“**Healthcare spending is outpacing what's financially sustainable in the long term**

¹ Source: Global Burden of Disease, Institute for Health Metrics and Evaluation (2022) | ² Source: WHO Health Workforce (2023) | ³ Source: OECD Health at a Glance (2019)

Now simpler, stronger and more focused



Fresenius Kabi

Fresenius Helios

Pharma

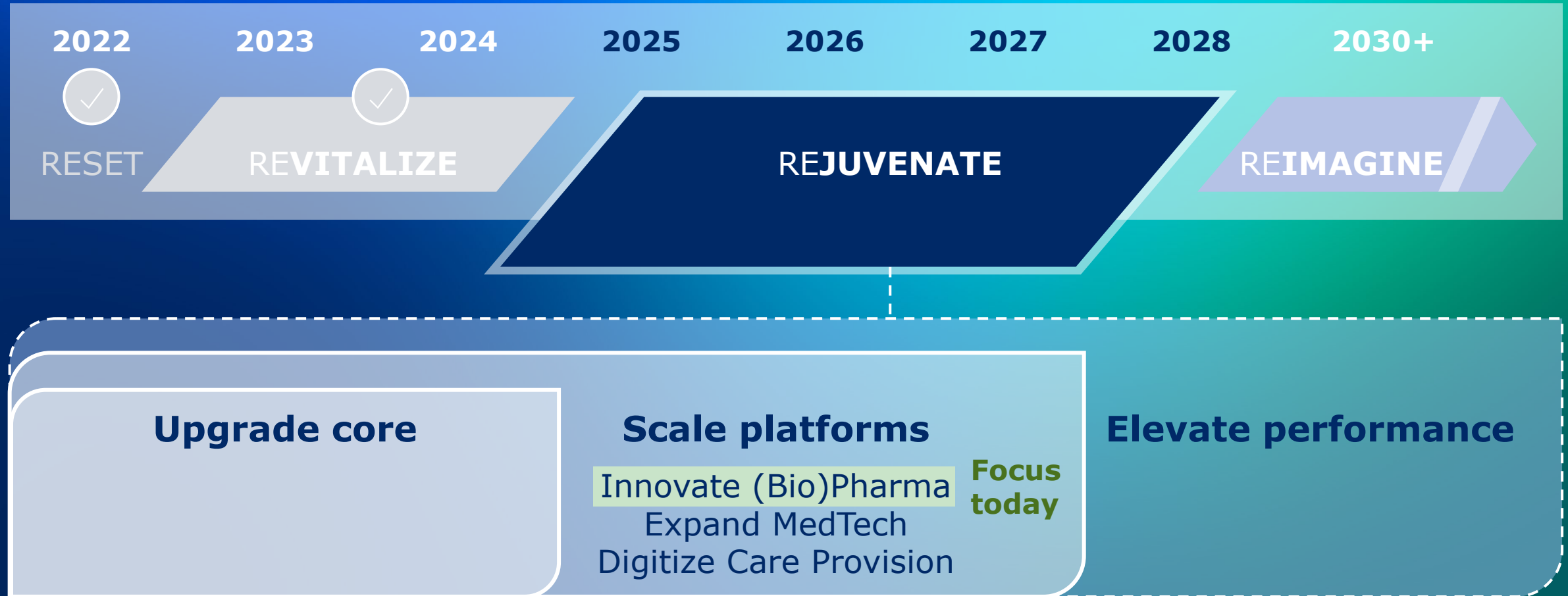
**Bio-
pharma**

Nutrition MedTech

Helios

Quirónsalud

Biopharma at the core of our Rejuvenate agenda



Biopharma is the next frontier for growth and patient access

As a leader in affordable and innovative healthcare products and highest quality patient care, Fresenius is expanding its biosimilars business to provide more patients access to biologic therapies

6x

Biosimilars market set to grow 6x by 2035 to >€180b¹ based on increasing number of upcoming LoEs

700m

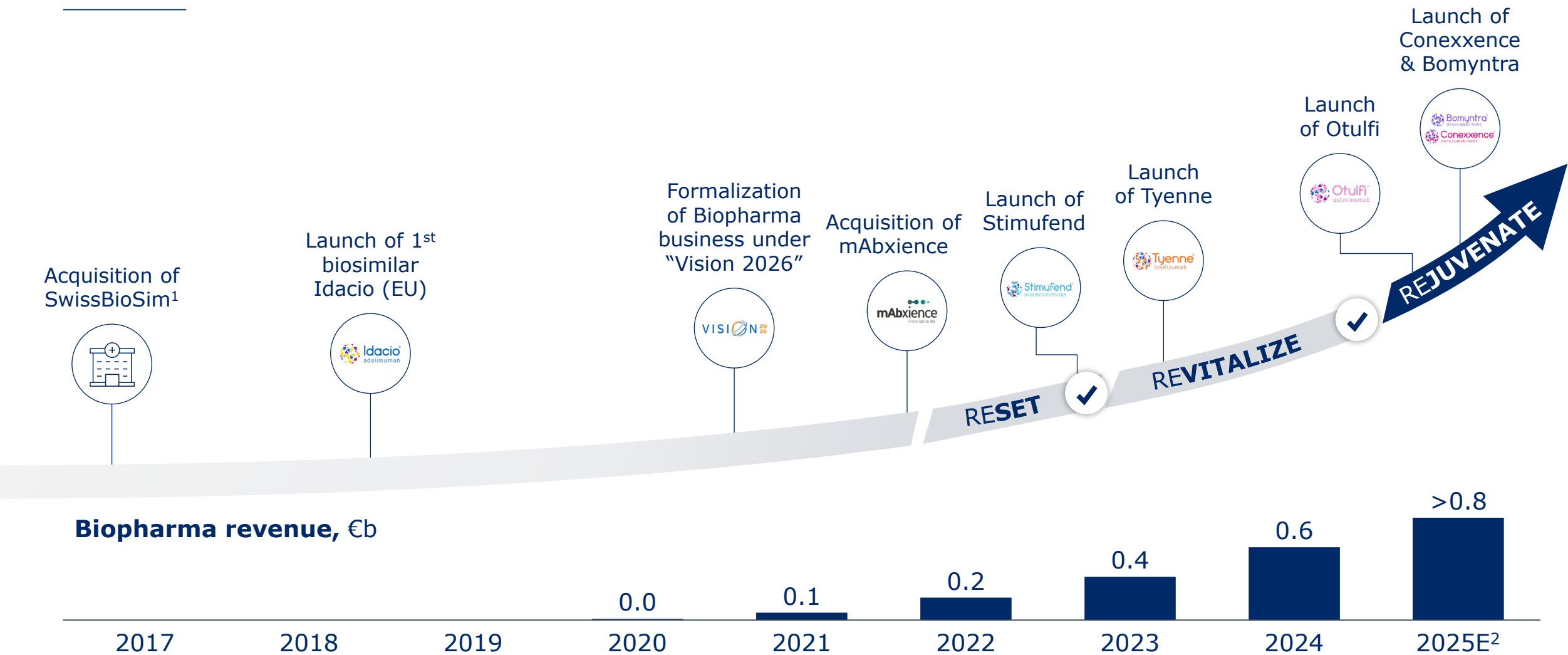
Patient days of biosimilar treatment globally² (2015-2023) prove broad adoption and patient access

€30b+

Annual savings across the EU and US – expected to grow to >€100b by 2030³

¹ Source: IQVIA, excluding GLP-1 molecules | ² Source: Biosimilars Council and US FDA | ³ Source: Biosimilars Council and IQVIA reports

Biopharma is a rapidly scaling business within Fresenius



¹ Merck KGaA created biosimilars business unit in 2011 | ² Vara consensus as of December 2025: ~€840m

Strong and renewed management team focused on execution



Leadership team with deep pharma and biotech expertise delivered successful scale-up

New governance and accountability structures implemented, revamped financial KPIs

Execution discipline with milestone-driven culture



Pierluigi Antonelli

President and CEO of Fresenius Kabi



Dr. Sang-Jin Pak

President Biopharma



Molly Benson

SVP Biosimilars US



Fabrice Romanet

SVP Research & Development



Michael Hammer

SVP Head of Portfolio & Business



Yannick Sorlet

SVP Head of TechOps, Supply Chain & Projects



Jurgen Van Broeck

CEO mAbxience



Laurent Rebier

SVP Partnerships



Niamh Furey

SVP Commercial EU & RoW Biopharma



Rachel Goode

SVP Legal and Intellectual Property



Michaela Rother

VP Global Head of HR



Sarah D'Orsie

SVP Global Government Affairs & Policy



Rolf Loesch

VP Compliance & Process Excellence



Tanja Greve

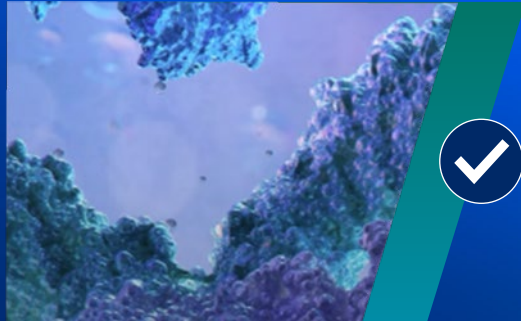
EVP CFO BU Biopharmaceutical



Catherine Priestley

VP Head of Communications Biopharma

#FutureFresenius: Rejuvenate in Biopharma



Biopharma is an attractive and rapidly scaling business within Fresenius



Fully vertically integrated Biopharma Powerhouse has the right levers to win in this highly attractive market



Capital allocation to drive further long-term profitable growth

II

Biopharma: value creation strategy

Our right to win

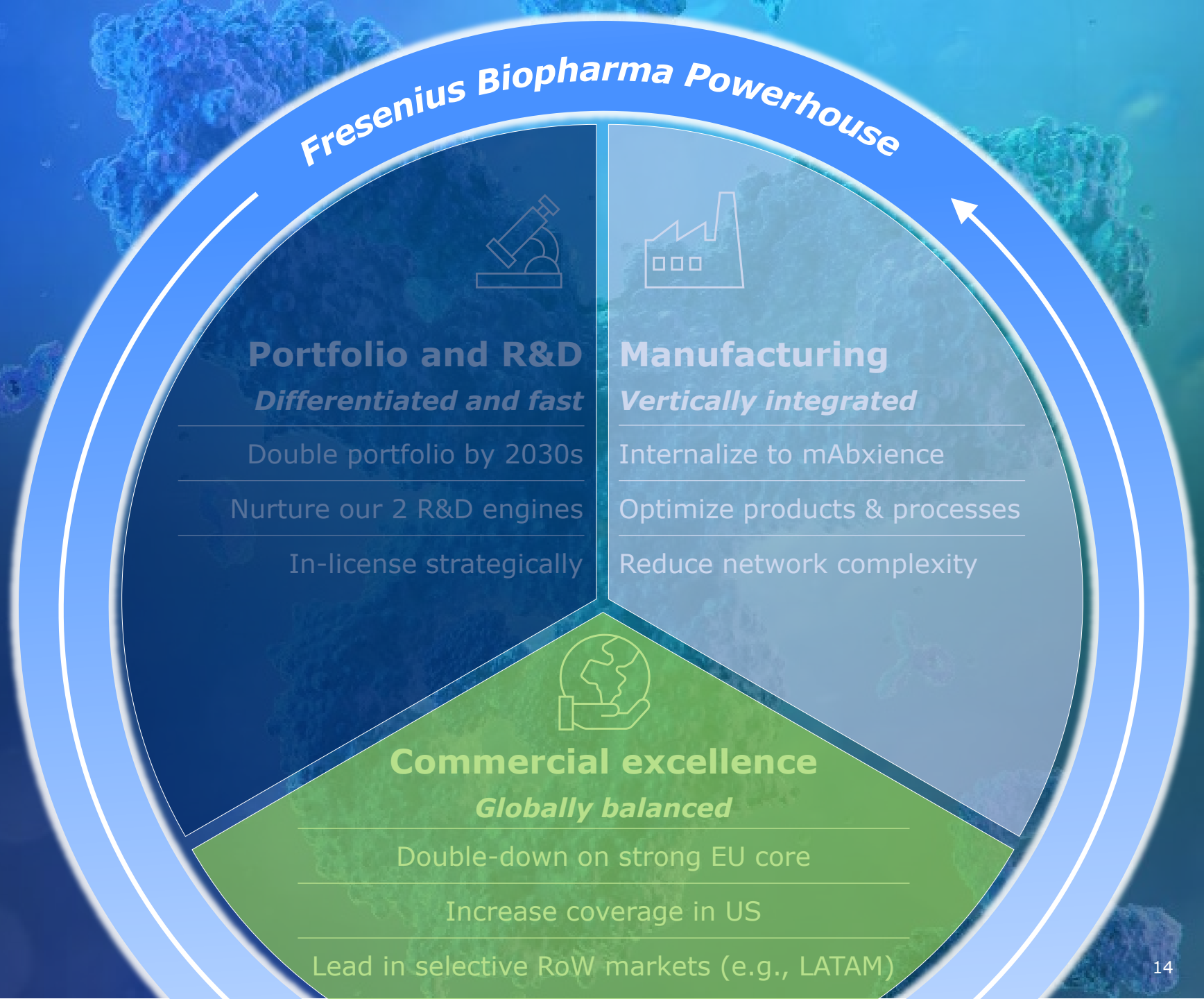


P. Antonelli



S.-J. Pak

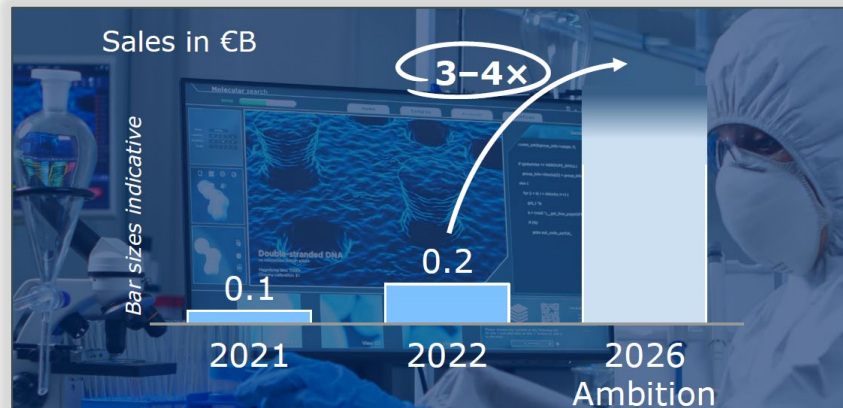
Biopharma: value creation strategy



Biopharma Powerhouse fully established – targets overachieved

Capital markets day 2023

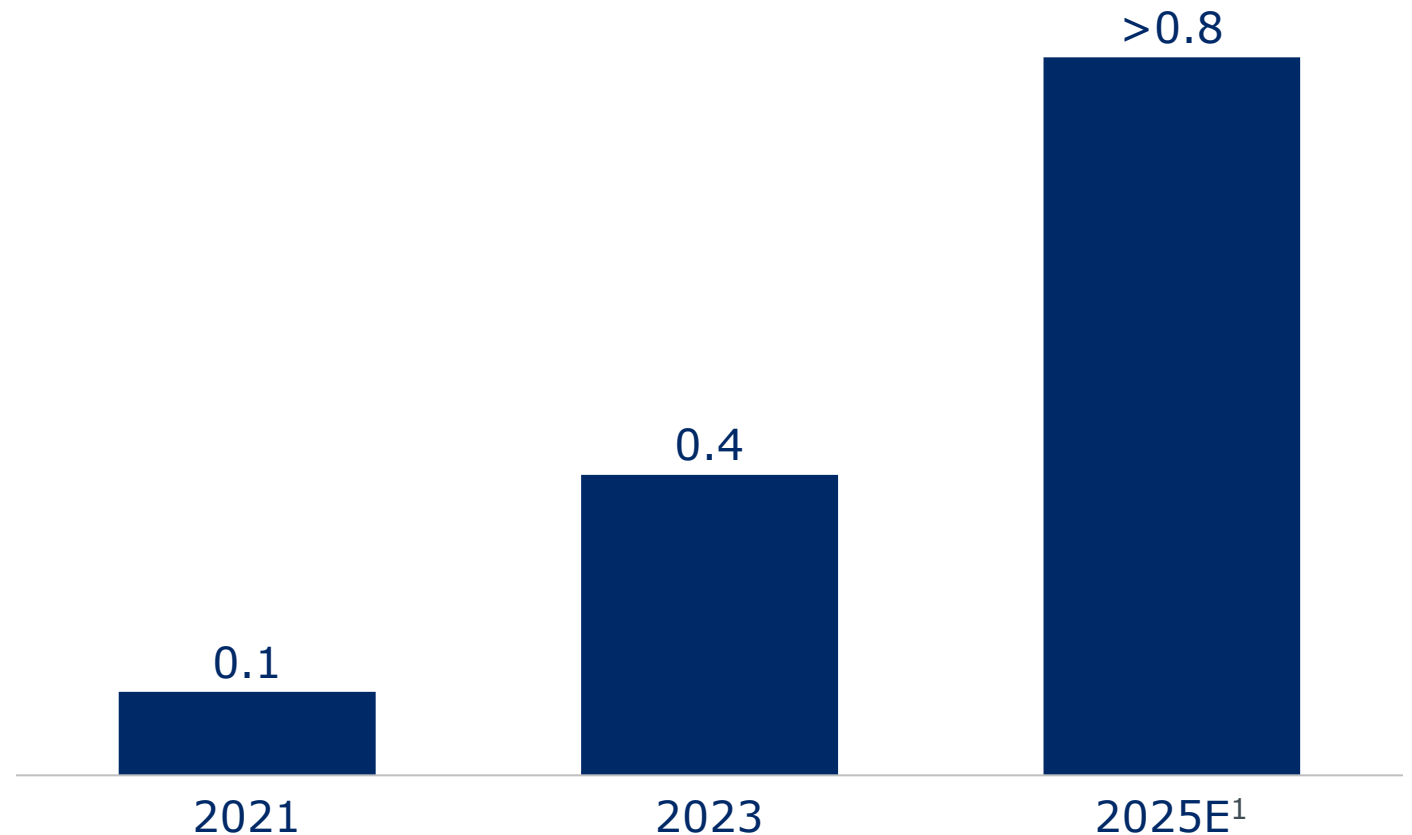
Biopharma revenue, in €b



Revenue ambition 2026
already reached in 2025E¹



EBITDA break-even target
overachieved in 2024

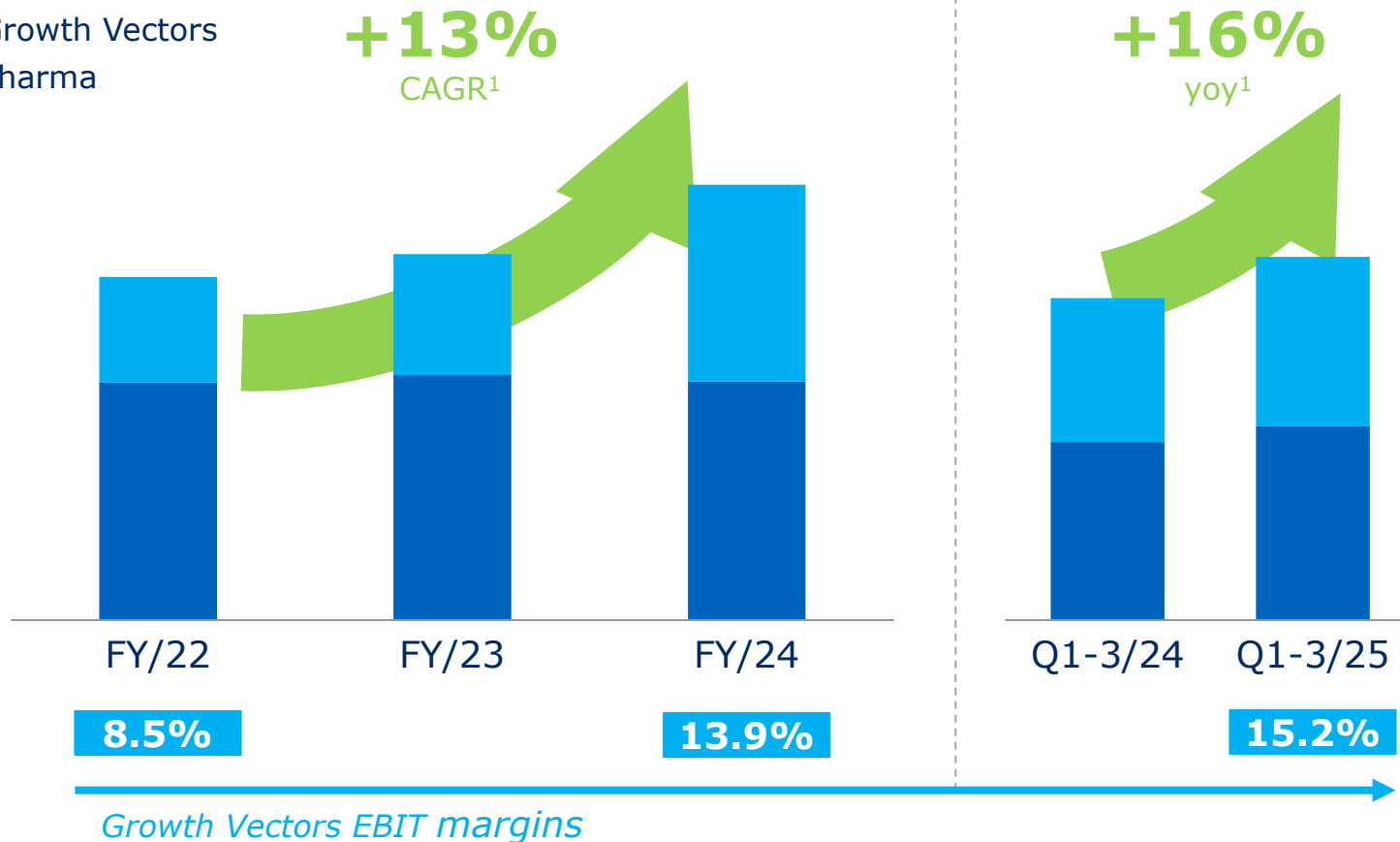


¹ Vara consensus as of December 2025: ~€840m

Biopharma is significantly contributing to EBIT growth

Fresenius Kabi EBIT

 Growth Vectors
 Pharma



Positive EBIT profile, step change in margins over recent years, moving towards Kabi's structural margin range²



Strong operating leverage as scale effects materialize



Pipeline ensuring **sustainable mid-term growth momentum**

¹ At constant currency and excluding Corporate | ² Margin band of 16-18%
Note: All figures before special items

Our right to win



Increasing Biopharma ambition for long-term profitable growth

~2x

Revenue^{1,2}



~20%

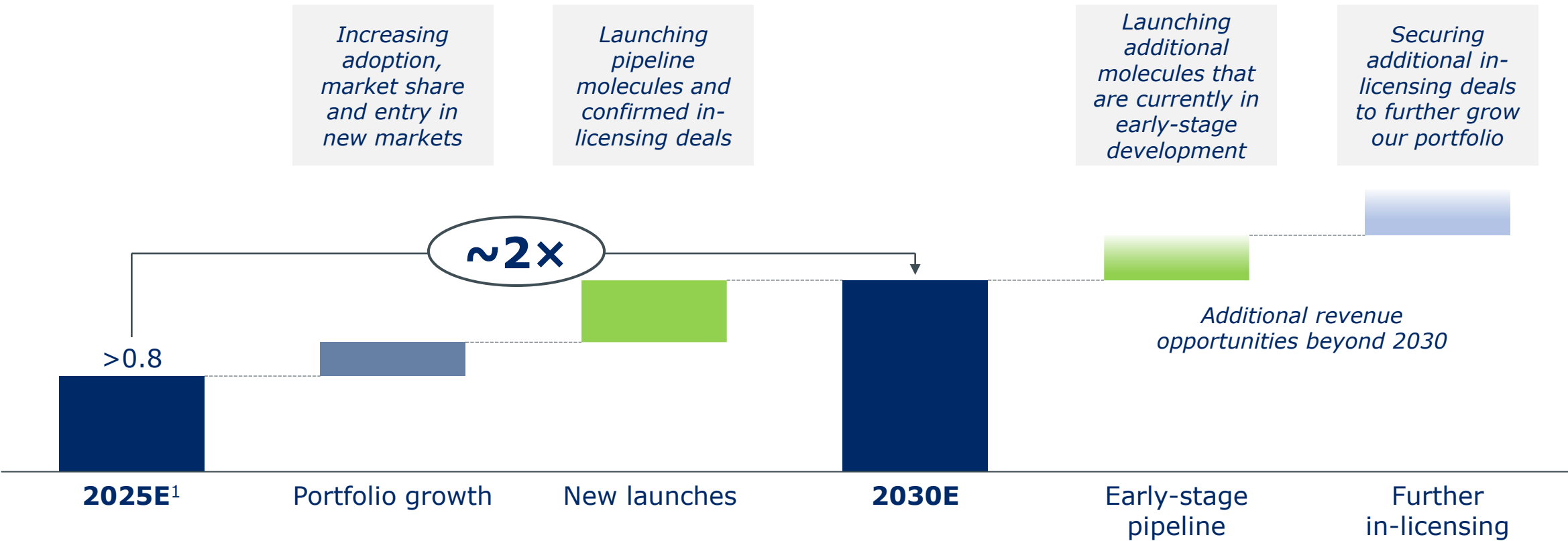
EBIT margin²

...by 2030

¹ Basis: FY/25 revenue; Vara consensus as of December 2025: ~€840m | ² Based on certain pipeline, portfolio, pricing, and market share assumptions

Poised to deliver upgraded ambition

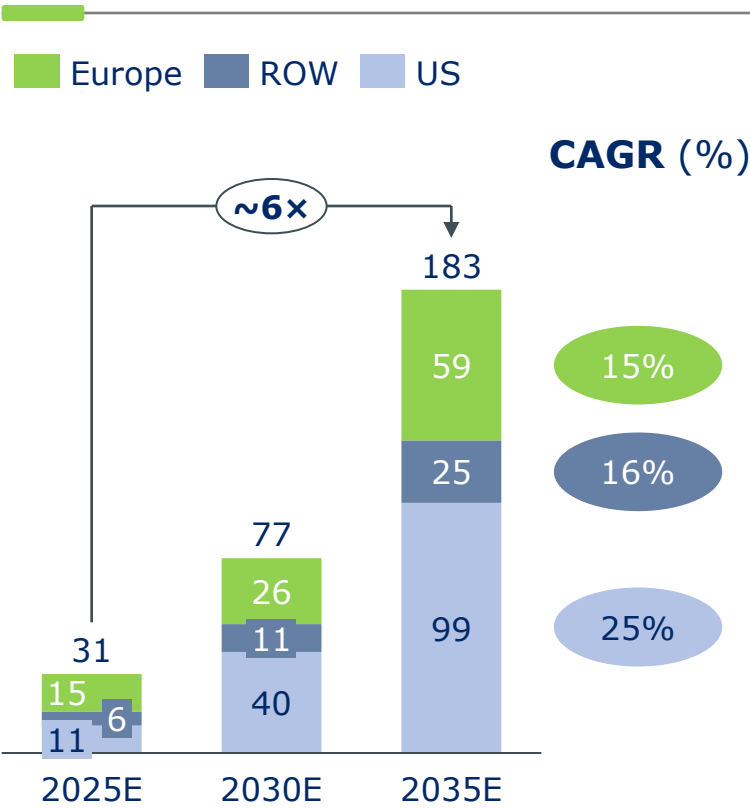
Revenue building blocks, in €b



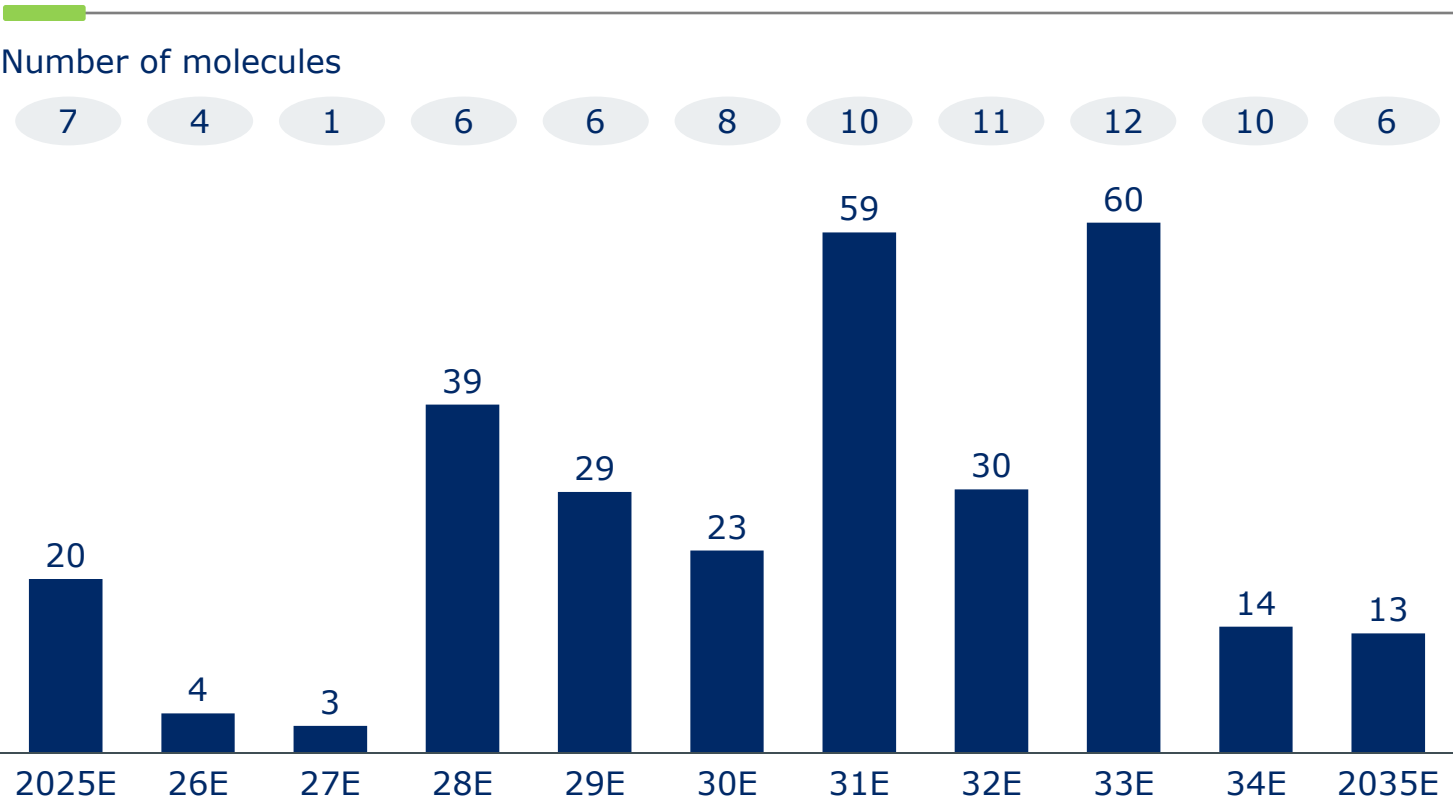
¹ Vara consensus as of December 2025: ~€840m
Note: Numbers are indicative

Global €31b biosimilar market set to grow 6x by 2035

Global biosimilar market¹, in €b



Worldwide originator sales facing Loss of Exclusivity, in LoE-1 sales per year² in €b

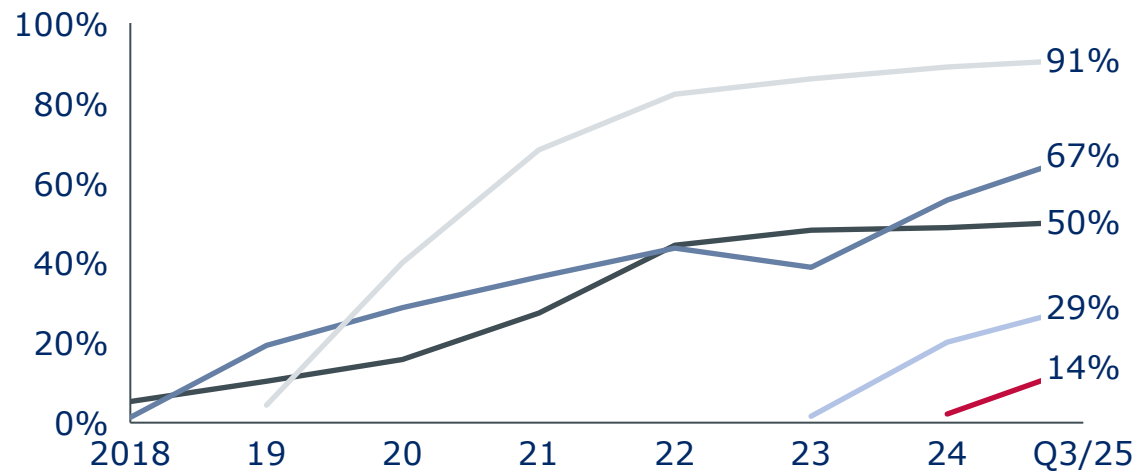


¹ Source: IQVIA. Excluding GLP-1 molecules. H2 2035E not available in IQVIA and forecasted based on CAGR 2030E - H1 2035E | ² Source: Evaluate Pharma. Evaluate Pharma considers the LoE year of the most critical patent (multiple patents with different LoE dates per molecule). Assuming Exchange Rate of €0.87 = \$1, including LoE-1 sales >€1b. Excluding GLP-1 molecules

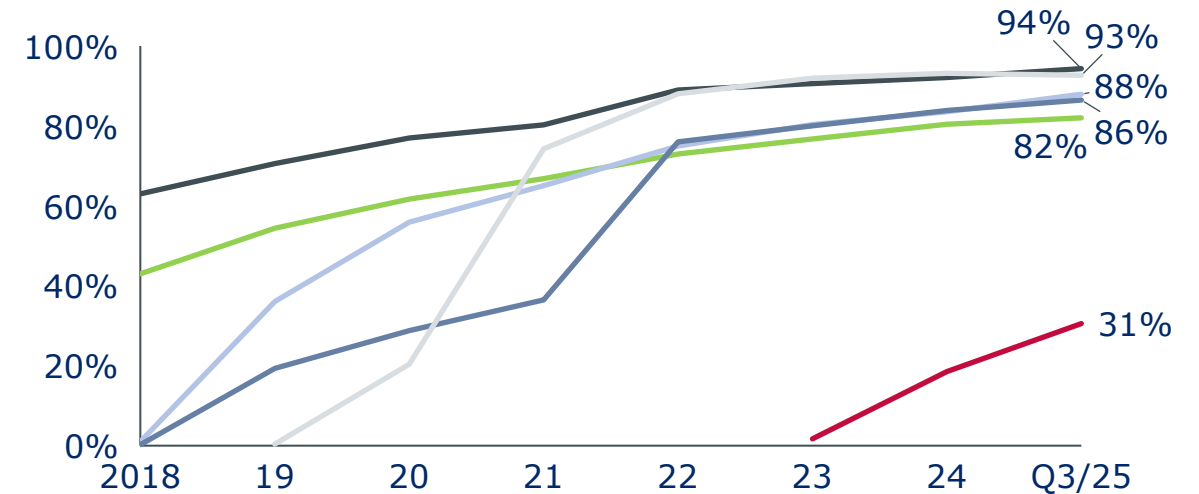
EU biosimilar adoption reaching >90% with US accelerating

Biosimilars share of total molecule volume¹, in %

US



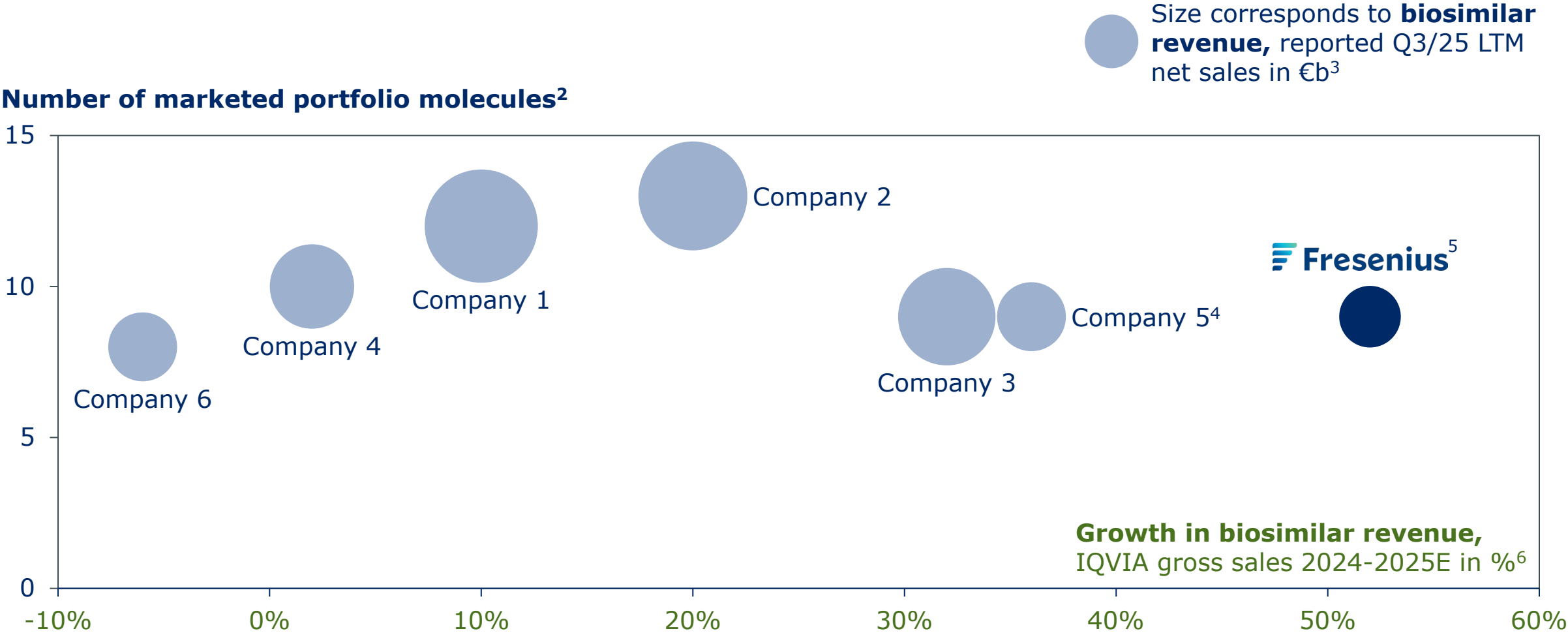
EU5²



As **familiarity** with biosimilars has **increased**, **adoption has accelerated** benefiting overall patient access and affordability

¹ Source: IQVIA – selective molecule examples | ² Includes Germany, UK, Italy, France, and Spain

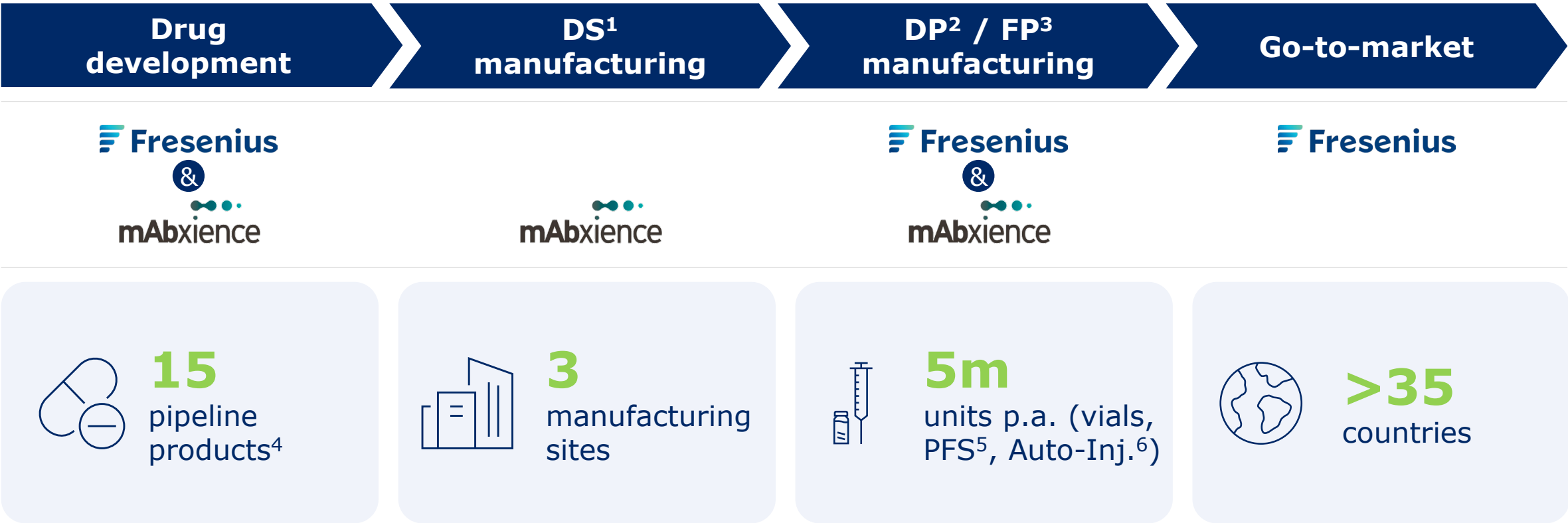
Fresenius is the fastest growing¹ biosimilar company globally



¹ Growth based on top-7 companies | ² Source: Company websites. Distinct molecules only - not counting multiple products per molecule | ³ Source: Company websites, assuming Exchange Rates of €0.87 = \$1, €0.0097 = ₹1, and €0.00059 = ¥1 | ⁴ Company 5 does not report biosimilar revenue, therefore IQVIA gross sales data is used | ⁵ Counting both Fresenius and mAbxience | ⁶ Source: IQVIA

Creating a leading end-to-end biosimilar business

Biopharma value chain



¹ Drug Substance (DS) | ² Drug Product (DP) | ³ Finished Product (FP) | ⁴ 15 products including 5 in registration/clinical, 4 in CMC/preclinical, 6 in early stage (6 not included in competitor comparison for consistency) |

⁵ Pre-filled syringe (PFS) | ⁶ Auto-Injector

Building a Powerhouse with broad portfolio of biosimilar brands

11 marketed products across 9 molecules



Idacio®
adalimumab



Stimufend®
pegfilgrastim-fpgk



Tyenne®
tocilizumab



Bomynta®
denosumab-bnht



Conexxence®
denosumab-bnht



Otulf®
ustekinumab

In-licensed



rituximab¹

Out-licensed



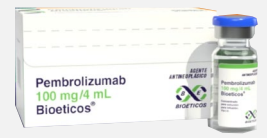
bevacizumab²

Out-licensed



pembrolizumab

Out-licensed



denosumab³
(osteoporosis)

Out-licensed

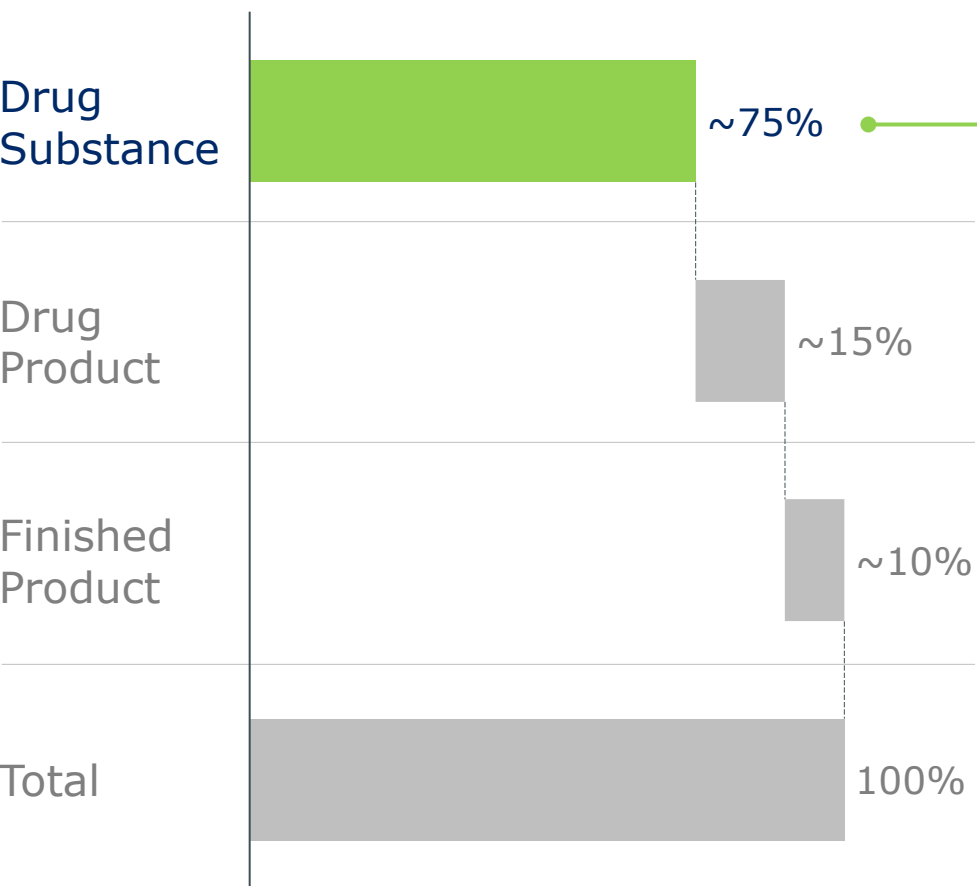
denosumab⁴
(oncology)

Out-licensed

¹ Referenced as Novex® in LATAM | ² Referenced as Alymsys®, among other names, in Europe, US and RoW | ³ Indicated for osteoporosis and referenced as Izamby® in Europe | ⁴ Indicated for oncology and referenced as Denbrayce® in Europe

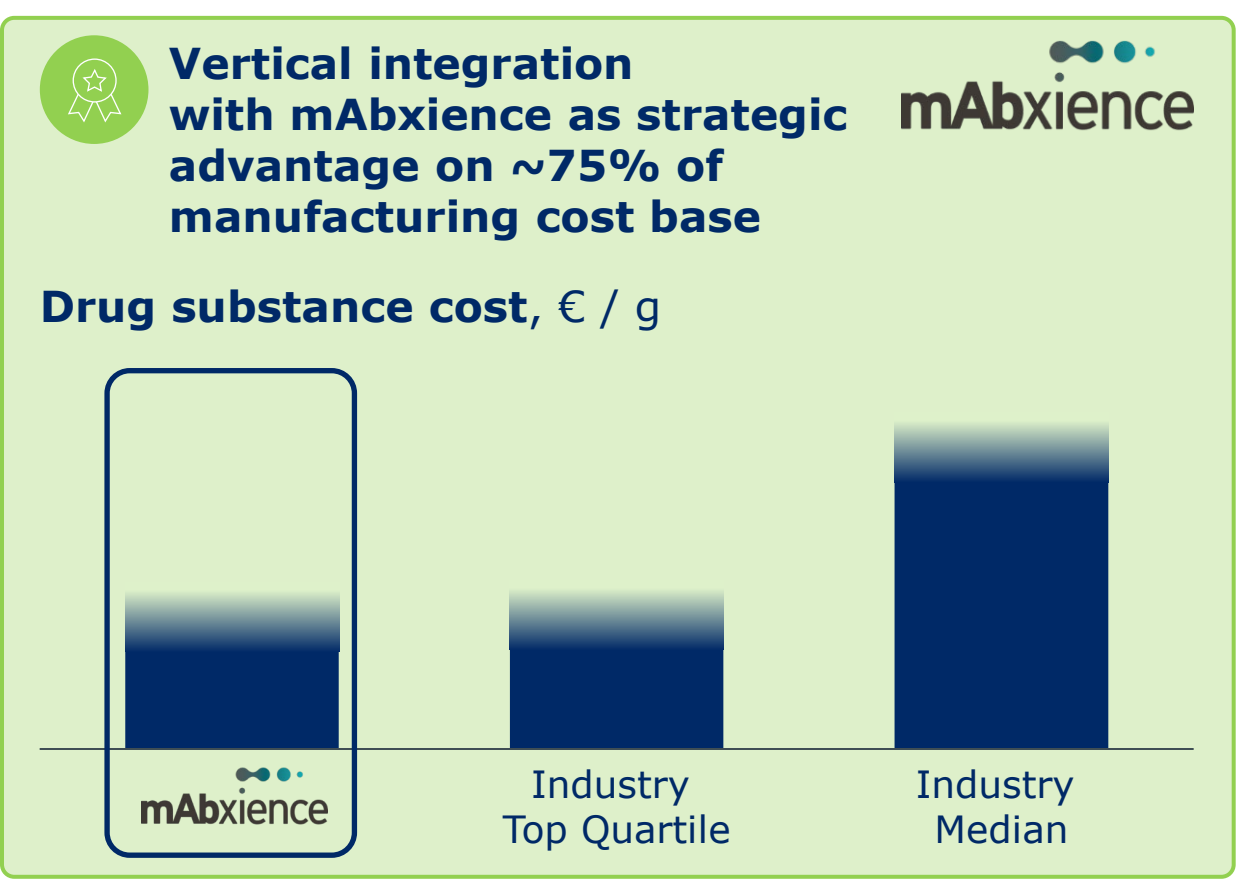
Vertical integration driving significant strategic cost advantages

Manufacturing cost¹



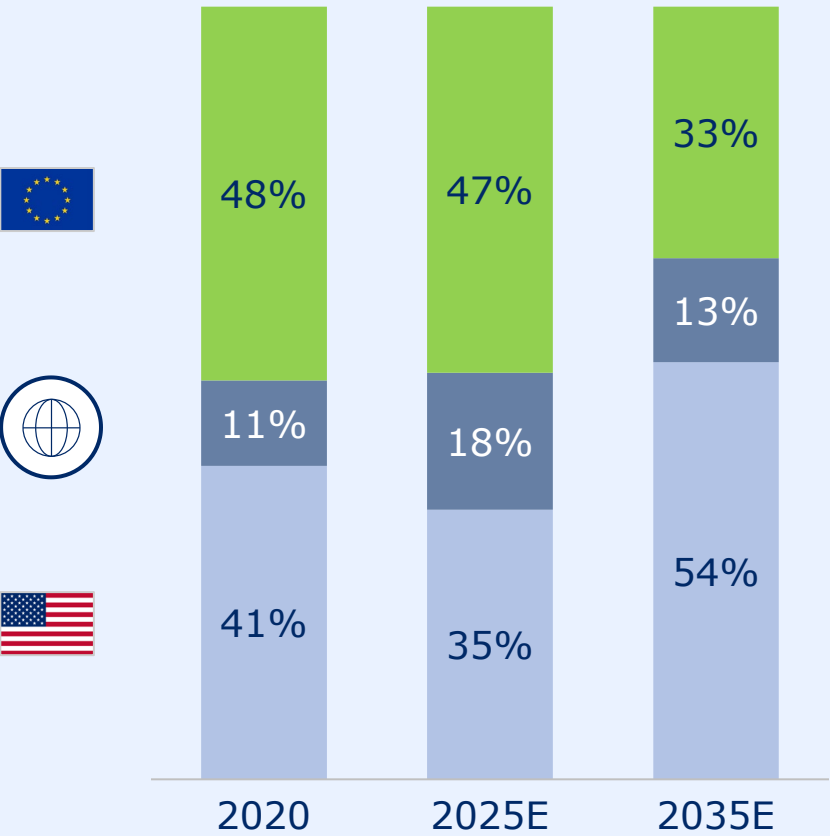
¹ Typical manufacturing cost split in Biosimilars industry

Our competitive advantage



Strong commercial presence across key geographies

Global biosimilar market¹



Our commercial coverage



Strong European core:

Direct-sales model in >20 markets and deep payer access (e.g., 95% tender win rates in France for tocilizumab, 100% of statutory sick funds contracted in Germany)



Leadership in LATAM:

Extensive commercial footprint across >10 countries and stronghold in Brazil and Argentina with favorable biosimilar regulation enabling early launches; ambition to opportunistically expand to selective MENA and APAC countries



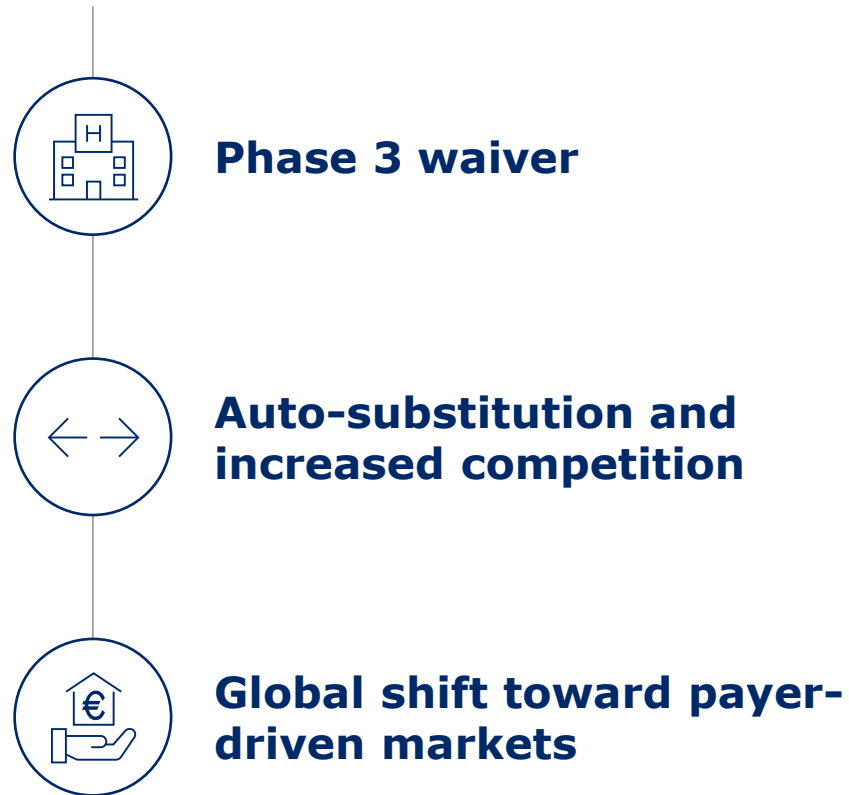
Ambition to increase US:

Continued portfolio expansion with recent US FDA approvals including novel commercial approaches to penetrate the market

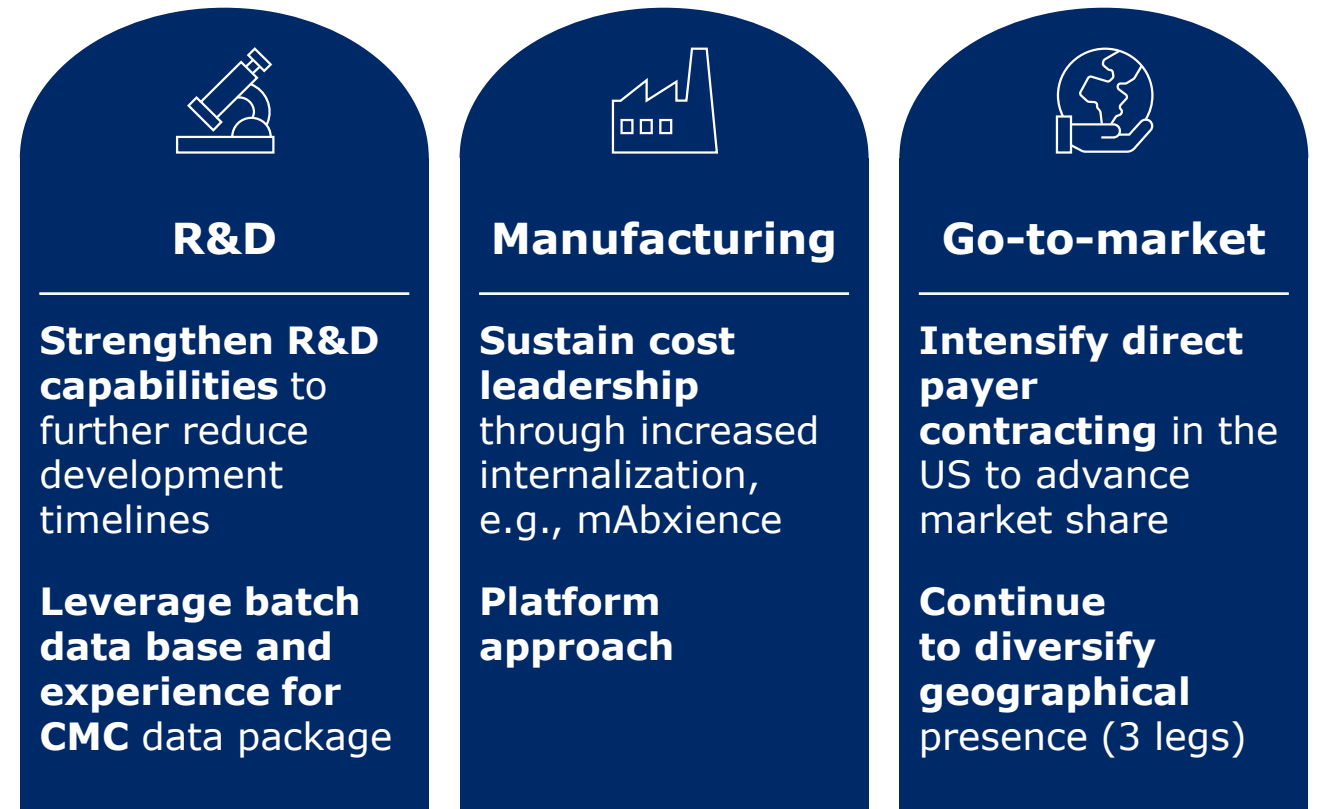
¹ Source: IQVIA. Excluding GLP-1 molecules. H2 2035E not available in IQVIA and forecasted based on CAGR 2030E - H1 2035E

Fresenius is positioned to address key shifts in biosimilar market

Market dynamics and shifts



Fresenius strategic response



Biopharma: value creation strategy



We operate in a dynamic market that is expected to grow sixfold over the next 10 years with increasing biosimilar adoption



We have established a global Powerhouse with a track record of successfully launching & scaling biosimilar brands over the last 6 years



We run an efficient R&D machine that delivers differentiated products at speed and continue to focus on selecting the right pipeline assets



We drive vertical integration incl. internalization of capacity, enabling cost leadership



We demonstrated the ability to penetrate key markets with tailored access strategies and geographic diversification

III

**Portfolio Rejuvenation
for long-term growth**

Our right to win

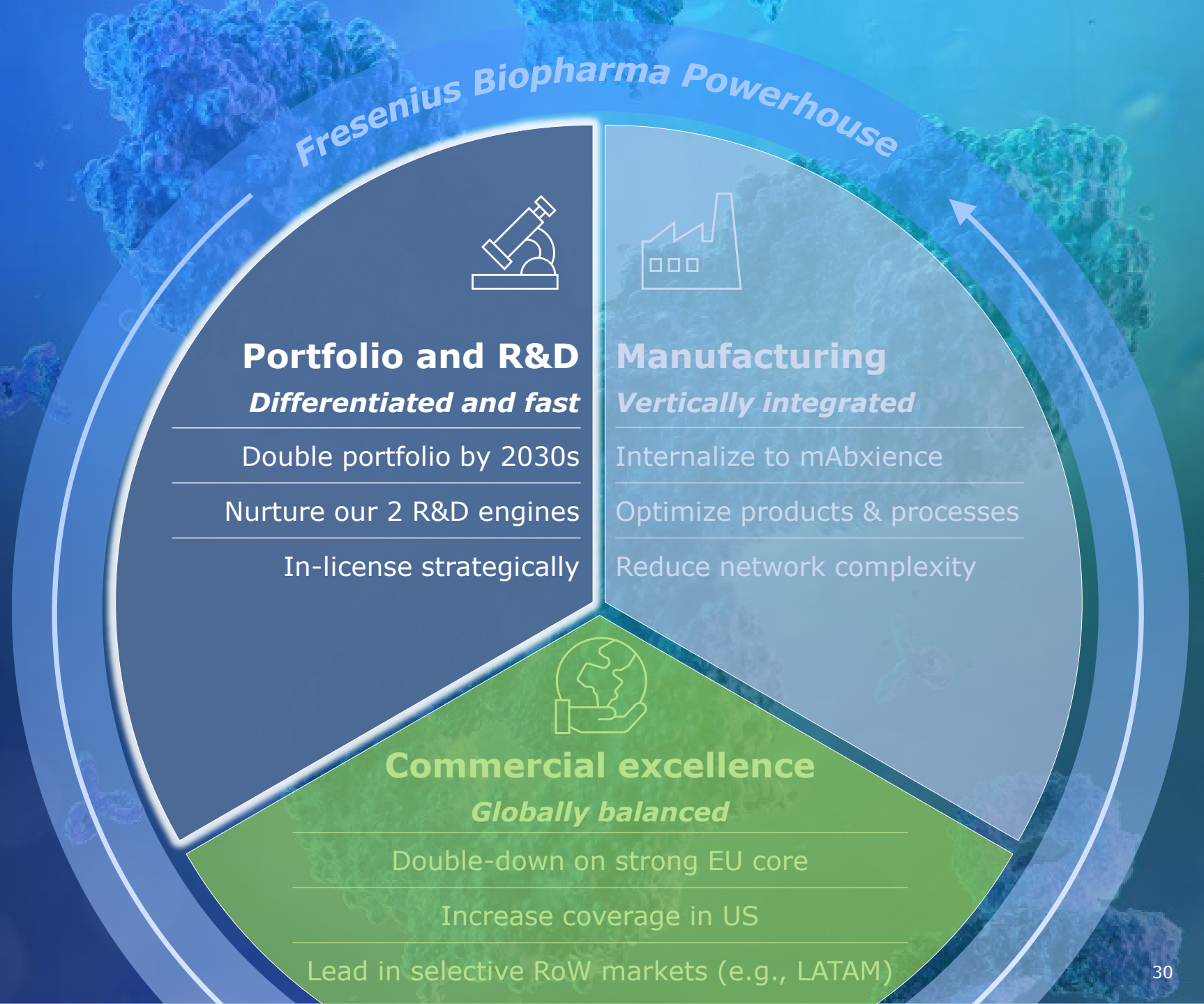


F. Romanet



M. Hammer

Portfolio Rejuvenation for long-term growth



Portfolio Rejuvenation for long-term growth



We have demonstrated both speed (e.g., tocilizumab first-to-market) and differentiation (e.g., denosumab pre-filled syringe) in R&D



We have built a competitive portfolio and pipeline covering €200b in originator sales



We will continue to deliver on our pipeline through our proven R&D engines, leveraging complementary in-house R&D hubs and selective in-licensing



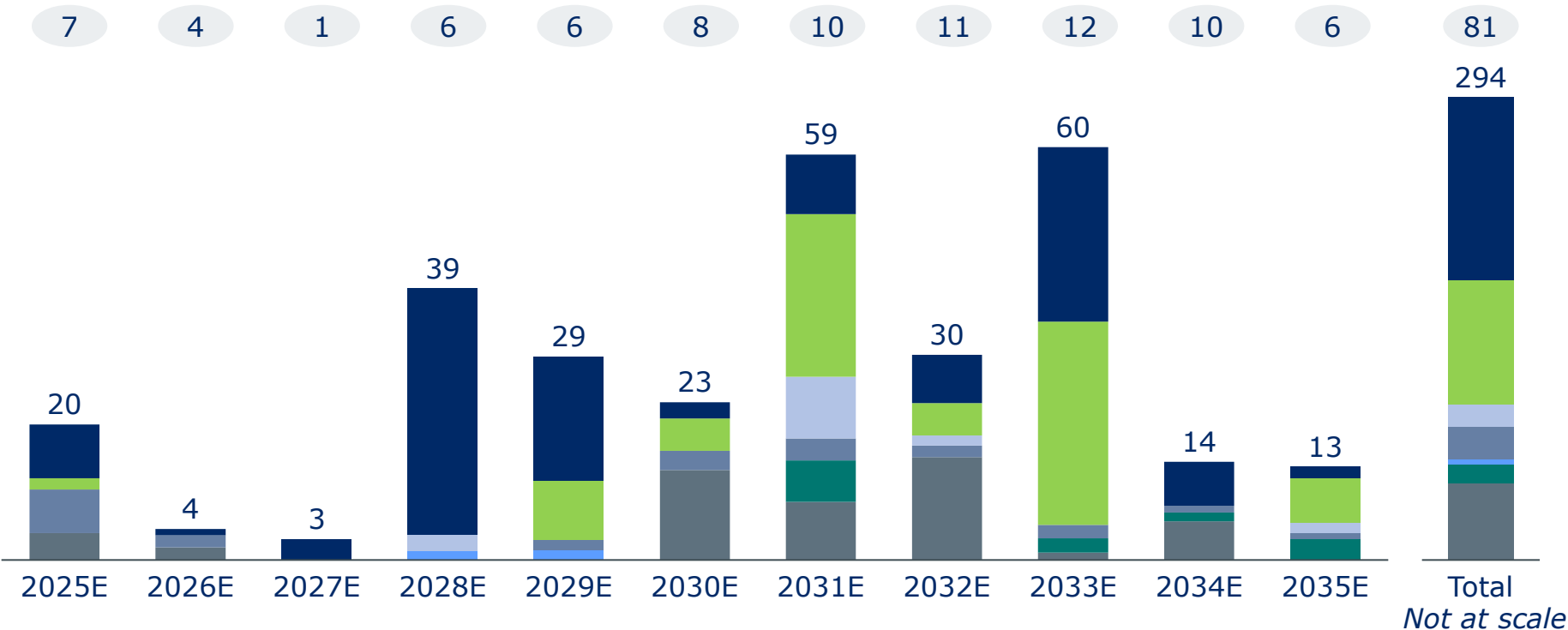
We target 2+ molecules p.a. entering development based on our competitive portfolio selection approach

Significant opportunity ahead from upcoming biologic LoEs

Therapeutic Areas (TAs)

■ Oncology ■ Immunology ■ Hematology ■ Musculoskeletal ■ Endocrinology (excl. GLP-1) ■ Neurology ■ Others ○ Number of molecules

Worldwide originator sales facing Loss of Exclusivity, in LoE-1 sales per year, in €b¹

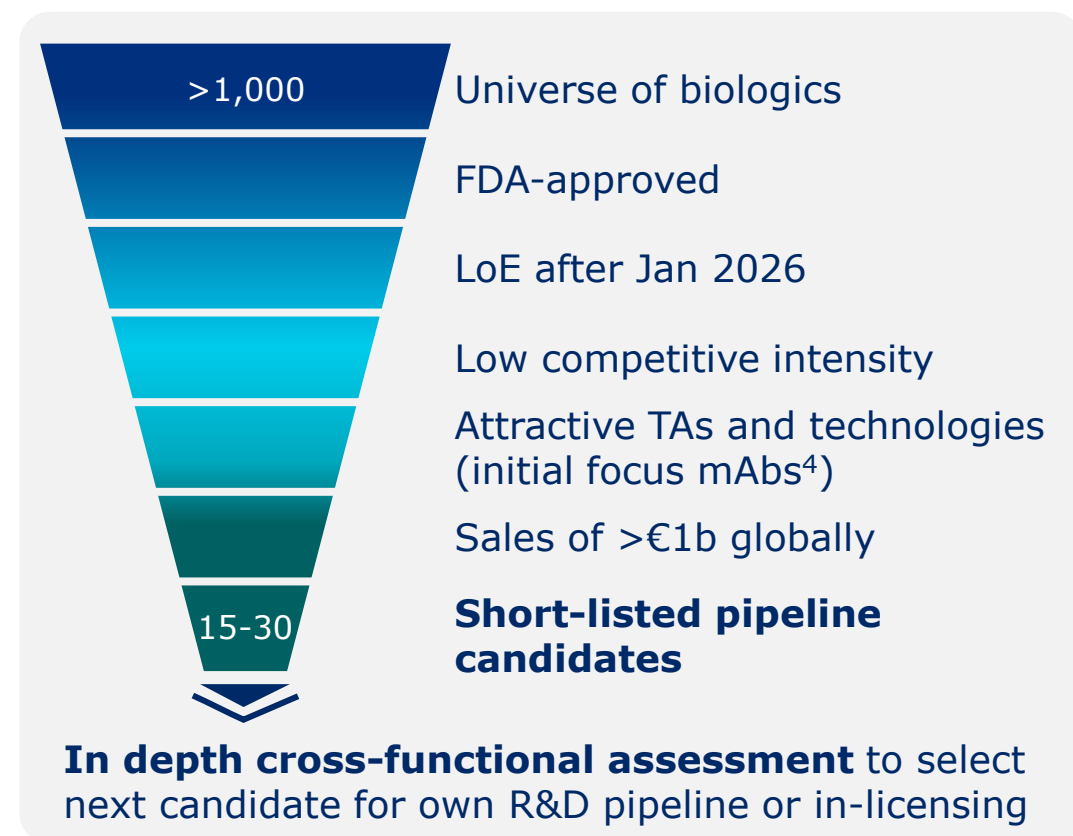


- > 60% of LoE-1 sales from Oncology and Immunology
- Experience in Oncology and Immunology enables strong positioning for upcoming launches
- Selective expansion into other attractive Therapeutic Areas

¹ Source: IQVIA. Assuming Exchange Rate of €0.87 = \$ 1, including LoE-1 sales > €1b. Excluding GLP-1 molecules

Targeted portfolio strategy and selection approach

Portfolio strategy and selection approach



Leveraging synergies between the immunology and oncology portfolio and **TA-agnostic expansion** through in-house development and partnerships



Strong portfolio execution prioritizing speed, best-in-class COP¹, full TPP², and reliable supply

2+

Molecules entering development p.a.

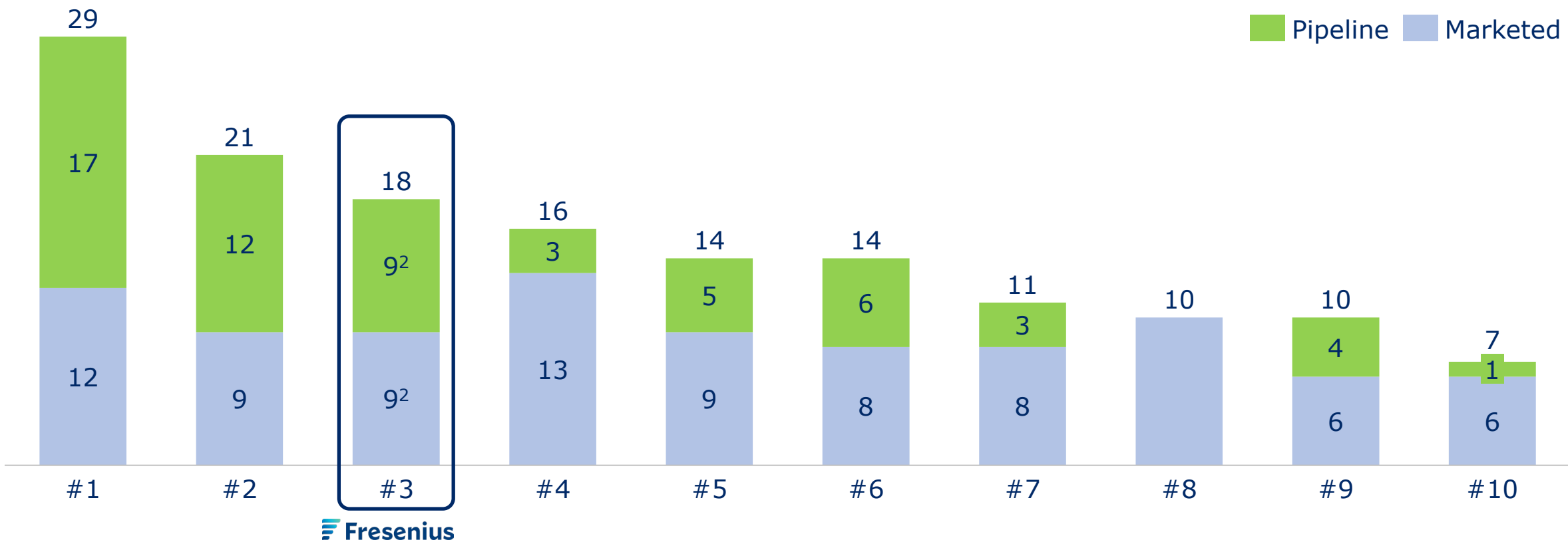
€200b

Portfolio & pipeline coverage of pre-LoE originator sales³

¹ Cost of Production | ² Target Product Profile | ³ Source: Evaluate Pharma | ⁴ Molecular Antibodies

Competitive portfolio across pipeline and marketed molecules

Number of portfolio molecules of top competitors globally¹ (excluding early-stage candidates)

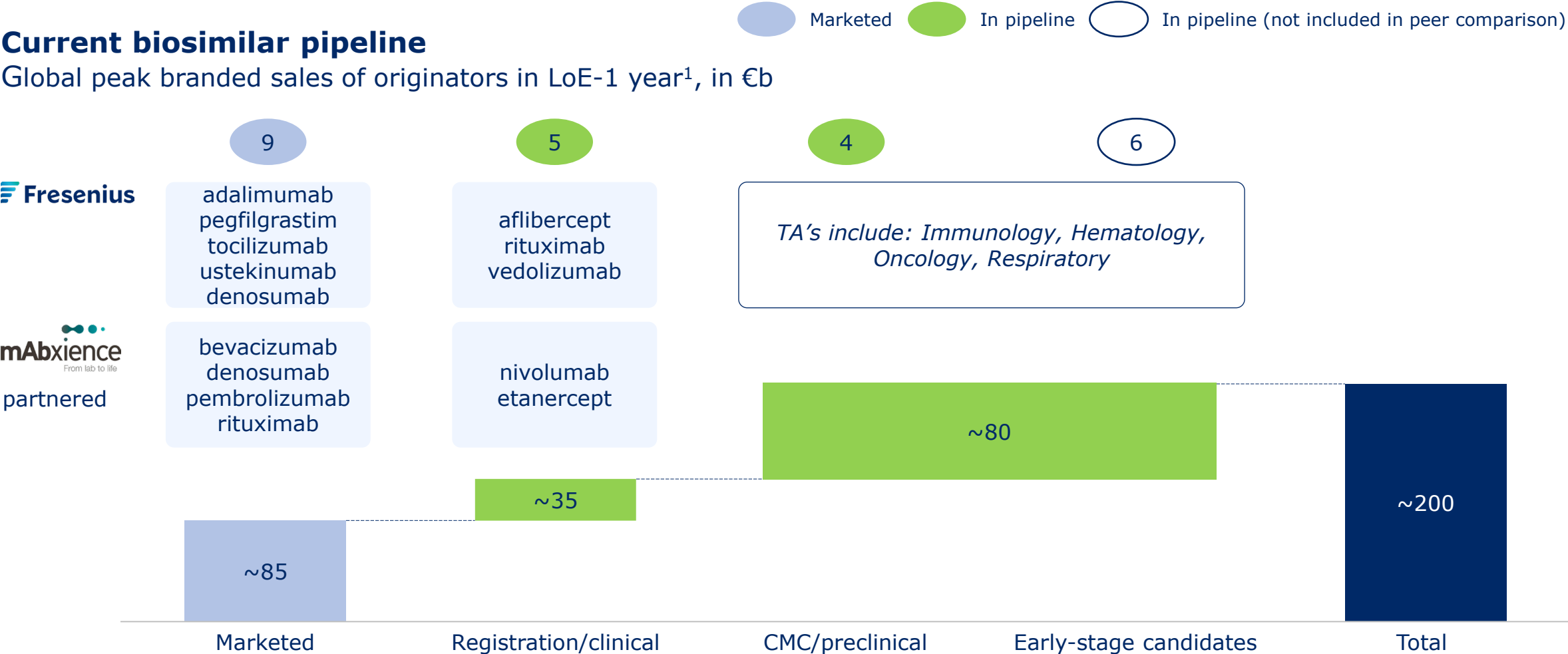


¹ Source: Company websites for pipelines from publicly disclosed information (December 2025). Disclosed pipelines only - actual pipelines might be larger. Distinct molecules only - not counting multiple products per molecule | ² Counting both Fresenius and mAbxience, excluding 6 early-stage candidates

Our portfolio & pipeline address €200b originator sales

Current biosimilar pipeline

Global peak branded sales of originators in LoE-1 year¹, in €b



¹ Source: Evaluate Pharma (accessed Jul 2025). LoE-1 year reflects timing of loss of exclusivity, sales figures may include molecules with LoEs pre-2025 where current market size has already declined

Complementary in-house R&D engines & strategic in-licensing

Complementary in-house R&D engines
increase portfolio breadth and competitive strength...



**...enhanced by
in-licensing**

 **Fresenius**
*Inhouse
R&D*


mAbxience
From lab to life
*Inhouse
R&D*

**Strategic
in-licensing**



Differentiation: Enabling launches with TPP differentiation



Speed: Enabling launches in first wave or first-to-market



Our in-house R&D engines enable increasing number of launches



Technical Development

15+ years of biosimilar development;
9 molecules launched,
9 in pipeline + 6 in early stage

Early-stage hubs in Eysins and León and **Scale-up hubs** in Garín, Munro, León



Clinical Development

Internal capabilities for pharmacokinetics, immunogenicity and biostatistics

Implementation of *estimands framework*¹ in all biosimilar study protocols since 2020



Regulatory & Quality

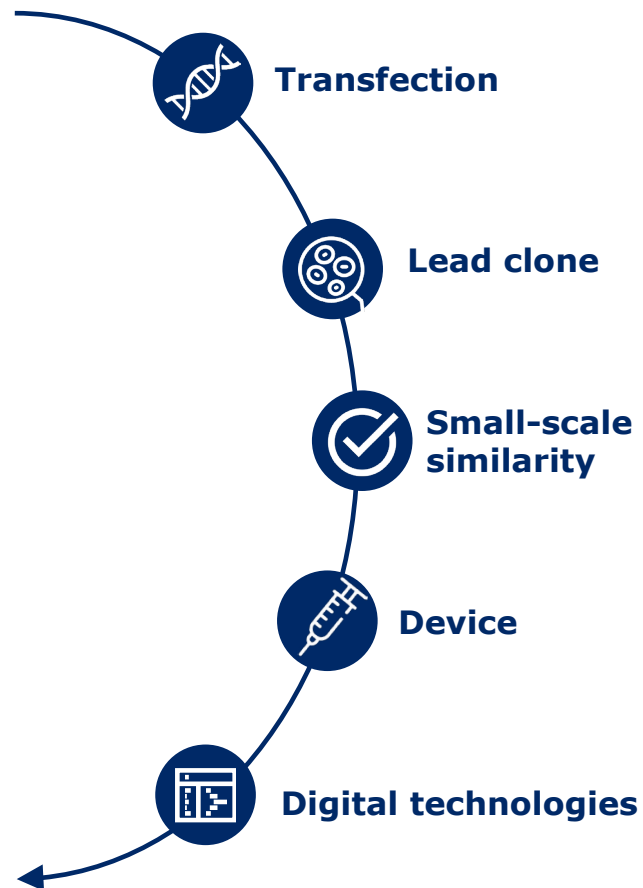
8 US FDA BLA approvals 2022-2025 – highest among competitors²

Leading voice in negotiating science-led adjustments of regulatory guidelines (e.g., BSUFA3³)

¹ An estimand is a precise description of the treatment effect reflecting the clinical question posed by the trial objective (as defined in ICH E9 R1) | ² Source: FDA BLA database | ³ Biosimilar User Fee Act 3

Our technical development allows differentiated & fast launches

Technical development steps



Shortening development cycle and ensuring high quality

- **Cell line development toolbox** proven to manufacture commercial products
- **State-of-the-art technologies** to pick the best producers and demonstrate clonality under industry-standard timelines
- **Combination of cell culture, media screening, and high-throughput analytics platforms** to select the lead clone, balancing quality, titer, and "clone plasticity"
- **Early establishment of the CMC blueprint** increases efficiency and accelerates timelines
- **Integrated CMC and IP experts** identify optimal manufacturing and formulation development
- **15 years of experience** allows to shorten development timelines with DOE and understand regulator expectation of right at first time approach
- **Platform approach for devices** reduces CDMO & internal development costs
- **Parallel development of multiple delivery technologies** enhances patient usability
- **Digital systems** (ELN, LIMS) ensure full data integrity and quality
- **Instrument connectivity and automated workflows** accelerate batch review & release

denosumab biosimilar is differentiated through pre-filled syringe

denosumab biosimilar overview of product differentiation

	 Fresenius	Originator	Competitor 1	Competitor 2	Competitor 3
Launch (US)	Jun 25		Jun 25	Jul 25	Nov 25
Dosage form					
Oncology PFS ¹	✓	✓			
Oncology vial	✓	✓	✓	✓	✓
Osteoporosis PFS ²	✓	✓	✓	✓	✓
Device specs					
	Latex-free	Latex-free	Latex-free	Natural rubber latex	Natural rubber latex

- 1 First-wave launch** securing early market entry and competitive positioning
- 2 Differentiated vs biosimilar competitors** through PFS¹ in oncology (e.g., solo bidder in UK tender in this category)
- 3 Prevents allergic reactions** through use of latex-free device

¹ 120mg | ² 60mg

Note: Additional competitors expected to launch in 2026

R&D speed showcased by first biosimilar launch of tocilizumab

Our US tocilizumab biosimilar launch



First-to-market launch through

- > Continuous regulatory dialogue
- > Parallelized clinical trials
- > Accelerated delivery of clinical study reports
- > Rolling publishing enabling fast delivery of final dossier

We in-license strategically to complement our internal pipeline

We are partner-of-choice

- ✓ **Deep expertise for development & regulatory**
- ✓ **Broad market access and trusted brand reputation**
- ✓ **IP & launch excellence**

Case Study: vedolizumab biosimilar in-licensing

Key information

Developer:  **polpharma**

Reference brand:  **Entyvio**

Therapeutic area: **Immunology**
Disease: **Inflammatory Bowel Disease (IBD)**

2024 global sales: **€5.8b**

Peak global sales: **€6.5b**



Fulfills program evaluation criteria

Portfolio fit

Therapeutic Area focus

Global agreement

Number of competitors

Program

Peak sales

First-to-market potential

Partner



EMA & FDA track record

Established supply chain

We shape and capitalize on the biosimilar regulatory landscape

Regulatory paradigm shifts

(incl. Phase 3 waivers)

Shorter **development timelines** and increasing **expectations for CMC analytical data package**

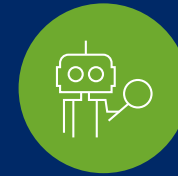
Increased need for **continuous FDA/EMA engagement**

Move towards **Phase 1 PK/PD focus**

Our competitive edge



40% faster development **timeline** achieved, and 15+ years of biosimilar expertise



10+ successful filings and AI-enabled regulatory tools



10+ robust **Phase 1 studies** with proven execution

Portfolio Rejuvenation for long-term growth



We have demonstrated both speed (e.g., tocilizumab first-to-market) and differentiation (e.g., denosumab pre-filled syringe) in R&D



We have built a competitive portfolio and pipeline covering €200b in originator sales



We will continue to deliver on our pipeline through our proven R&D engines, leveraging complementary in-house R&D hubs and selective in-licensing



We target 2+ molecules p.a. entering development based on our competitive portfolio selection approach

IV

**Cost Leadership:
reaping benefits of vertical integration**

Our right to win



Y. Sorlet



J. Van Broeck

**Cost Leadership:
reaping benefits of
vertical integration**



Cost Leadership: reaping benefits of vertical integration



We operate a cost-leading manufacturing Powerhouse across Drug Substance, Drug Product, Finished Product - with global certifications



We achieved significant COGS reduction through internalization and will continue to leverage our full vertical integration



We target additional COGS reduction through product and process optimization initiatives on capacity, productivity, and supply chain efficiency



With our end-to-end manufacturing network established, we are investing >€300m over the next 5 years to expand capacity and drive growth

Our industry-leading manufacturing capacity & capabilities

Global manufacturing footprint

León 		24 kl Bioreactor cell culture capacity	✓ 2 industrial lines and cGMP¹ pilot facility ✓ Single-use technology
Garín 		24 kl Bioreactor cell culture capacity	✓ 2 independent industrial lines ✓ Single-use technology
Munro 		400 l Bacterial fermentation capacity	✓ Dedicated process optimization lab ✓ Single-use technology
Graz 		5 m units Finished product capacity p.a.	✓ Semi-automated assembly of devices ✓ Single-use and isolator technology

¹ Current Good Manufacturing Practice

We continue to invest in expanding our capacity

CapEx
investment
of >€300m
over the
next 5
years



León

Increased bioreactor size and additional production line: DS in new building, doubling capacity



Munro

New microbiological site: Capability added to manufacturing network



Graz

Additional production line: Compounding & filling for vials and syringes, more than doubling capacity



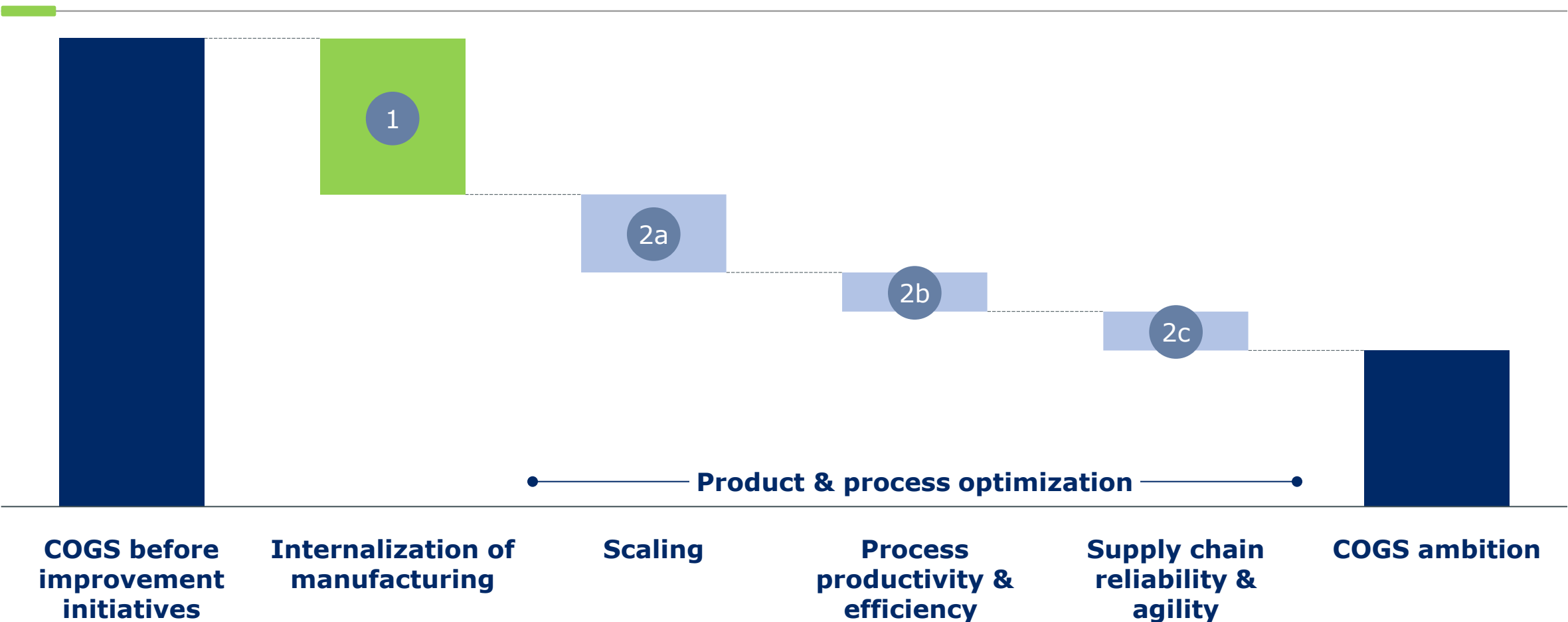
Graz

Additional production line: Automated packaging for Auto-injector/PFS, more than doubling capacity and reducing costs

¹ Additional investment in additional capacity/technology not included in CapEx number above

Improving COGS through internalization & process optimization

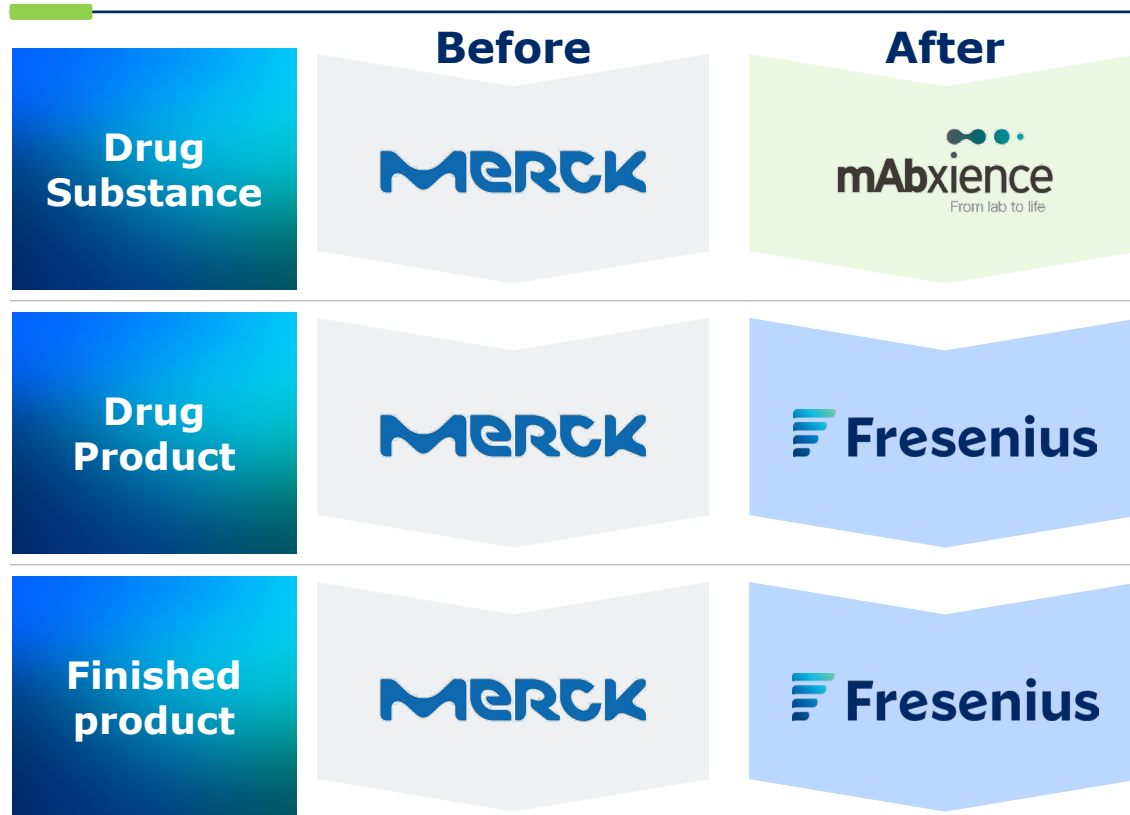
COGS reduction initiatives (indicative)



1 Internalization

We are on track to further insource to our internal sites

Supply chain internalization (example tocilizumab biosimilar)



Internal manufacturing capacity coverage (across portfolio)

Internal DS, in % of total capacity required



Internal DP, in % of total capacity required



Internal FP, in % of total capacity required

>80% of capacity already in 2025

1 Internalization

mAbxience as our core platform for internal DS manufacturing

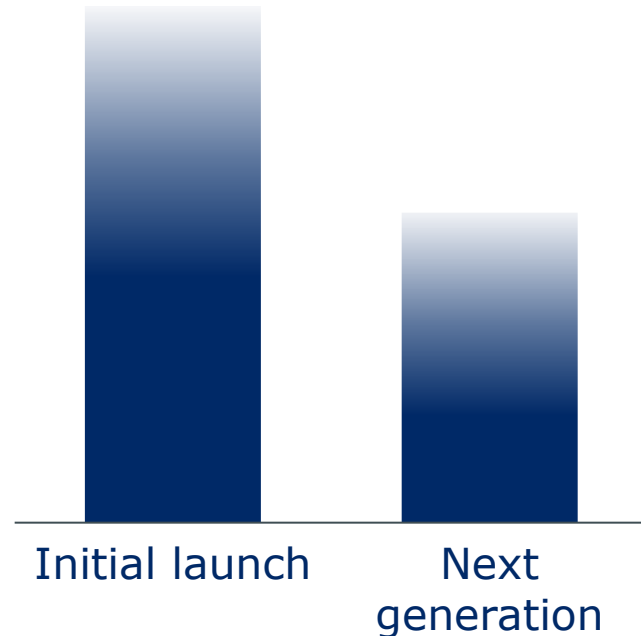
Integration on track, with additional opportunities for further improvement



We continuously improve productivity even within same scale

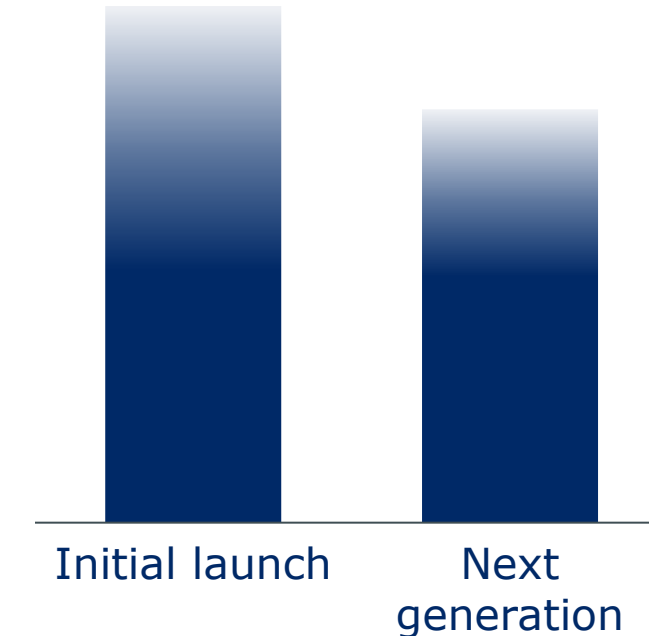
DS cost reduction bevacizumab biosimilar

Drug substance costs



DS cost reduction tocilizumab biosimilar

Drug substance costs



Improvements of productivity and costs in each process iteration – even within same scale

- > Enhancement of cell productivity
- > Increase in yield
- > Reduction of raw material costs

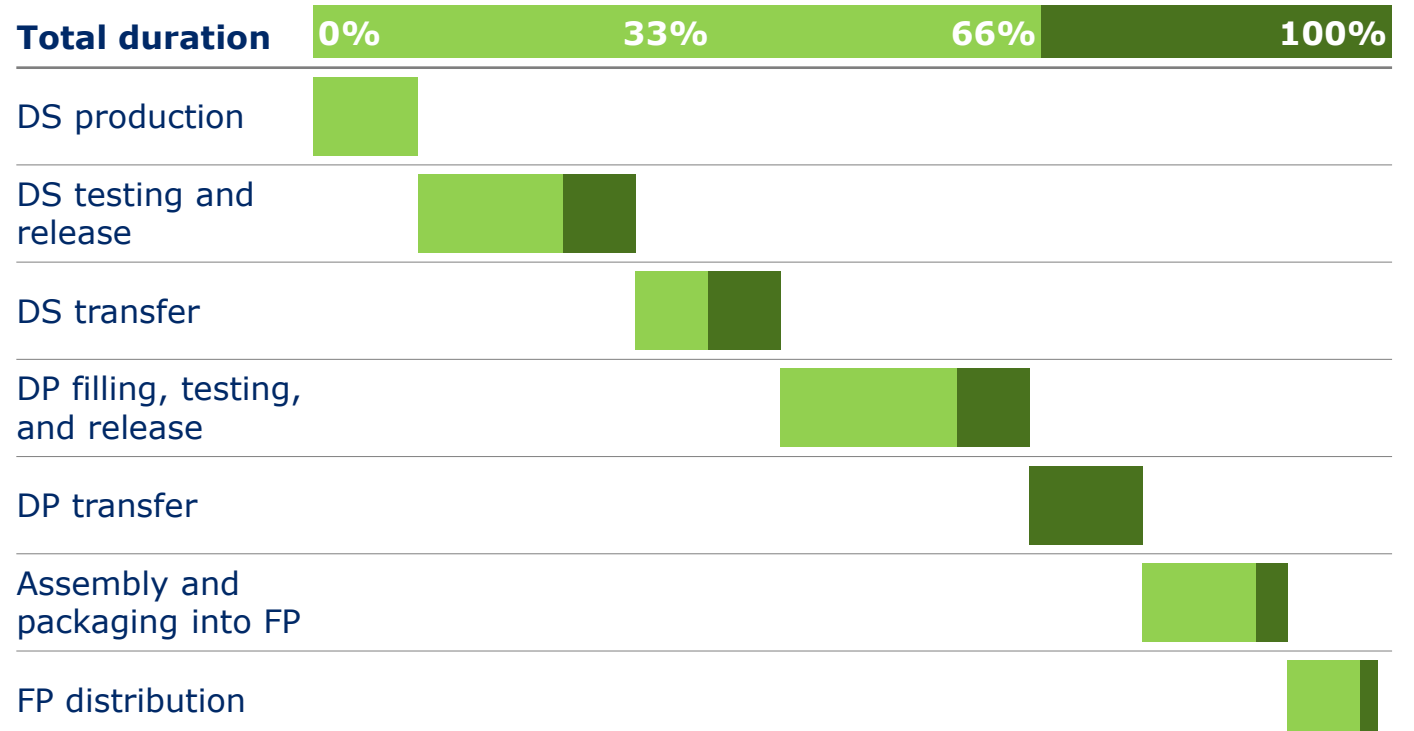
Relentless execution to reduce lead time by 33%

Increased supply chain agility

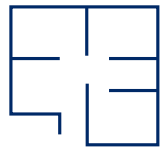
- > Manufacturing products closer to market demand
- > Reducing cycle stocks, inventory levels and value, as well as inventory risks
- > Shortening the end-to-end production timeline

Lean manufacturing (lead time reduction by 33%)

Duration of processing steps (indicative): ■ Ambition ■ Reduction



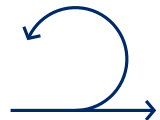
We will be at the forefront for future manufacturing



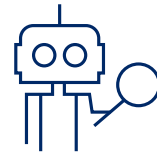
Flexible capacity model (e.g., different scales)



Geographic footprint expansion for DS/DP capacity



Manufacturing technology, e.g., continuous manufacturing



Automation & digital manufacturing, e.g., for sampling, material flow, packaging

Cost Leadership: reaping benefits of vertical integration



We operate a cost-leading manufacturing Powerhouse across Drug Substance, Drug Product, Finished Product - with global certifications



We achieved significant COGS reduction through internalization and will continue to leverage our full vertical integration



We target additional COGS reduction through product and process optimization initiatives on capacity, productivity, and supply chain efficiency



With our end-to-end manufacturing network established, we are investing >€300m over the next 5 years to expand capacity and drive growth



**Commercial excellence:
balanced footprint across geographies**

Our right to win



S.-J. Pak



M. Benson

**Commercial
excellence:
balanced footprint
across geographies**



Commercial excellence: balanced footprint across geographies



We execute successfully across key regions: deep payer access in EU, gaining traction in US, leader in LATAM, scaling access in other regions



We have a targeted go-to-market strategy molecule by molecule and have shown first-to-market ability with Tyenne, the leading tocilizumab biosimilar



We will continue to de-risk our revenue streams through our commercial network and selected partners with milestone payments

Continue targeted mix of direct sales & out-licensing partners

Revenue split by channel (indicative)

2023



2025E



2030E

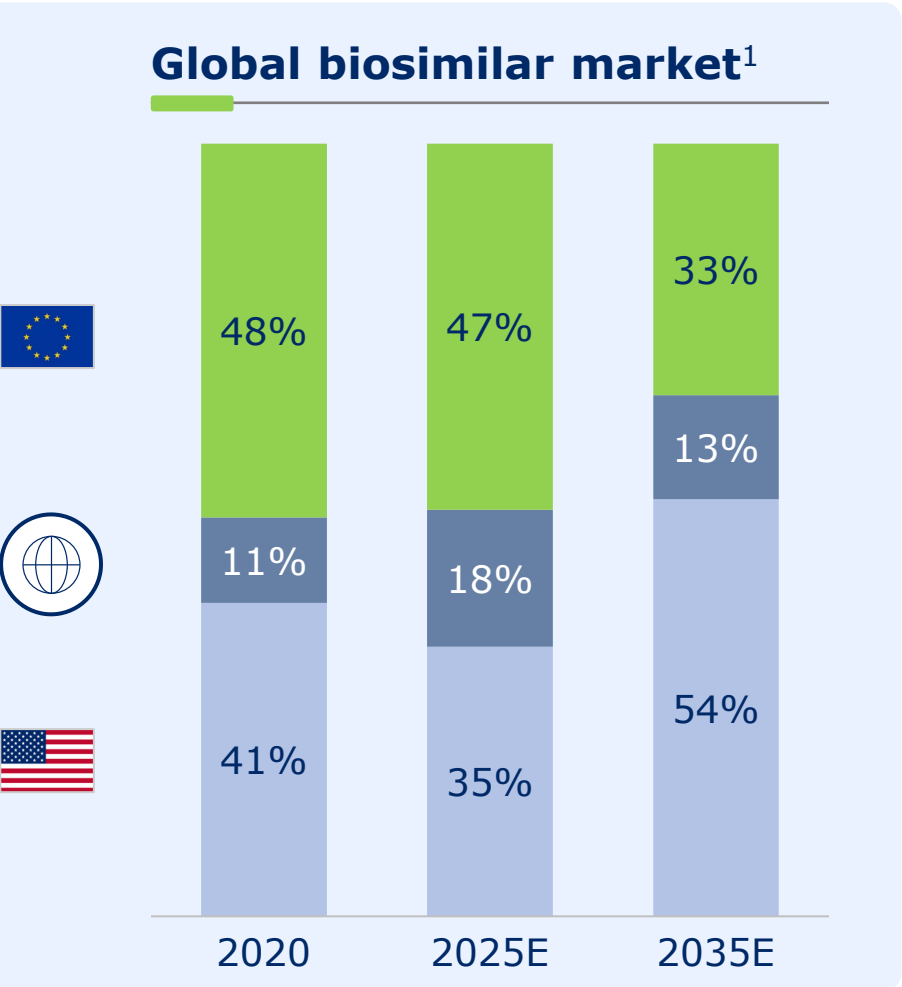


■ Fresenius direct sales ■ mAbxience out-licensing partnerships

As our commercial network expands, **we will increasingly generate revenues through our own sales engine** – raising margins and reducing dependence on milestone payments

We will continue to leverage out-licensing partnerships selectively and strategically

Strong commercial presence across key geographies



Our commercial coverage



Strong European core:

Direct-sales model in >20 markets and deep payer access (e.g., 95% tender win rates in France for tocilizumab, 100% of statutory sick funds contracted in Germany)



Leadership in LATAM:

Extensive commercial footprint across >10 countries and stronghold in Brazil and Argentina with favorable biosimilar regulation enabling early launches; ambition to opportunistically expand to selective MENA and APAC countries



Ambition to increase US:

Continued portfolio expansion with recent FDA approvals and use of novel approaches to penetrate the market

¹ Source: IQVIA. Excluding GLP-1 molecules. H2 2035E not available in IQVIA and forecasted based on CAGR 2030E - H1 2035E

Spotlight EU: Building a scalable, integrated commercial model

Well established direct presence

- ✓ Direct presence in >20 European markets in both public and private provider systems
- ✓ Experienced regional key account management with direct engagement model with hospital procurement & clinical stakeholders

25+ leading hospitals and hospital purchasing groups for Tocilizumab



Competitive Edge in Tenders

- ✓ Pan-European tender framework harmonizing tender management system
- ✓ Expertise in multi-country, multi-channel bidding

40+ tenders won in 2025

Strong performance management

- ✓ Analytics solution combining, e.g., external sales data, customer facing team metrics
- ✓ Optimization of resource allocation, messaging, and actions

32% market share in tocilizumab biosimilars

Spotlight LATAM: Fresenius leads the adalimumab market

Capabilities demonstrated in Brazil



- ✓ **Incubator market** for piloting ahead of global launch (e.g., adalimumab biosimilar)
- ✓ **#1 biopharma market** in LATAM
- ✓ **#1 market position** based on >40 years of Fresenius presence
- ✓ **Productive Development Partnership (PDP)** with 10-year supply agreement and 40% minimum public market share

Capabilities demonstrated in Argentina

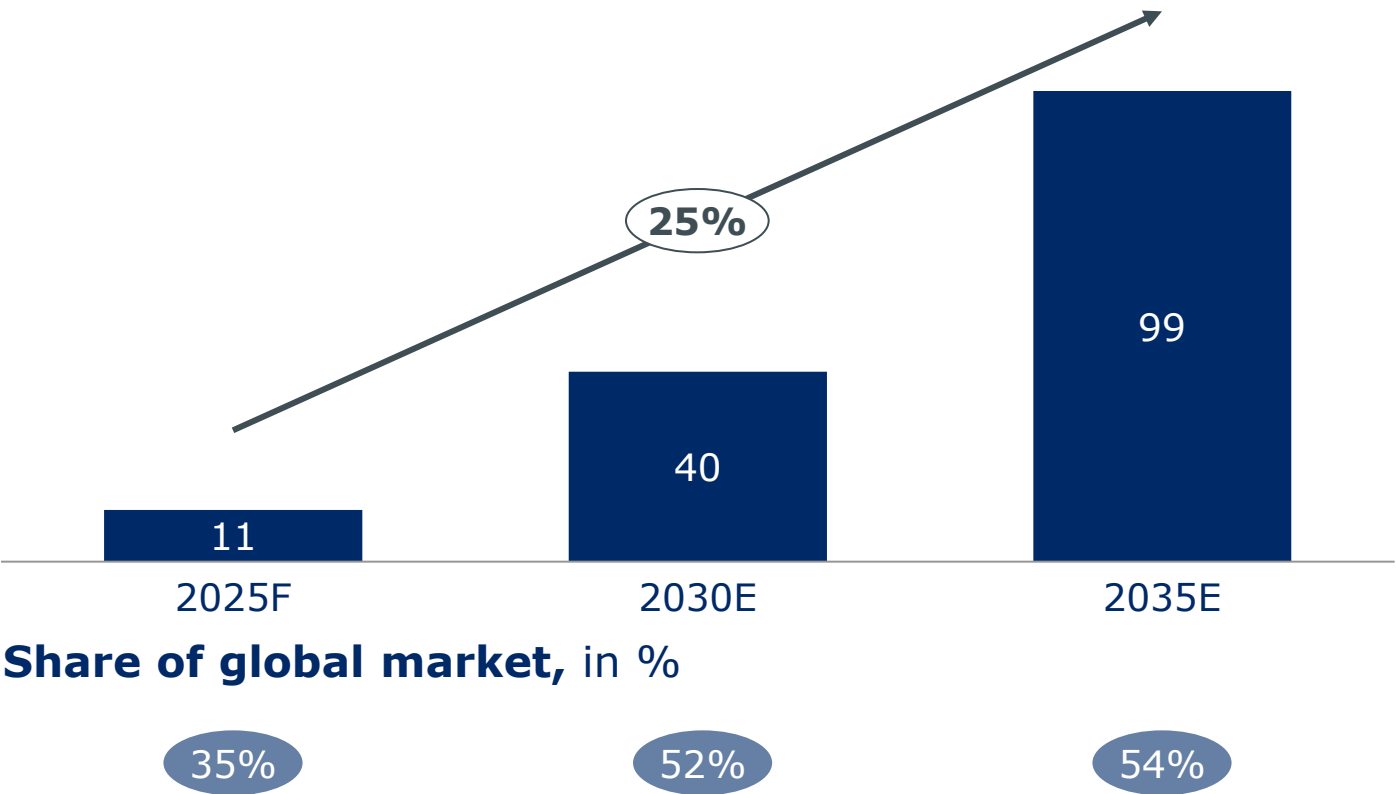


- ✓ **Incubator market** for piloting ahead of global launch (e.g., pembrolizumab biosimilar)
- ✓ **Significant biopharma market** in LATAM
- ✓ **Strong local market position** of our products
- ✓ **Dual national-provincial tender structure** informed our robust tender capabilities that are relevant for other tender-driven markets



Spotlight US: Attractive, high-growth, and evolving market

US biosimilar market¹, in €b



Market evolutions

- > **IRA² legislation opens** early-entry windows for authorized biosimilars
- > **Independent and employer health plans** shift toward transparent pricing outside traditional PBMs³
- > **Increasing interchangeability designations** accelerate biosimilar uptake

>**60%** Revenue growth 24-25E

¹ Source: IQVIA. Excluding GLP-1 molecules. H2 2035E not available in IQVIA and forecasted based on CAGR 2030E - H1 2035E | ² Inflation Reduction Act. Introduces drug pricing reforms that enable earlier competition with originator products | ³ Pharmacy Benefit Manager



Tyenne is the first and leading tocilizumab biosimilar in the US



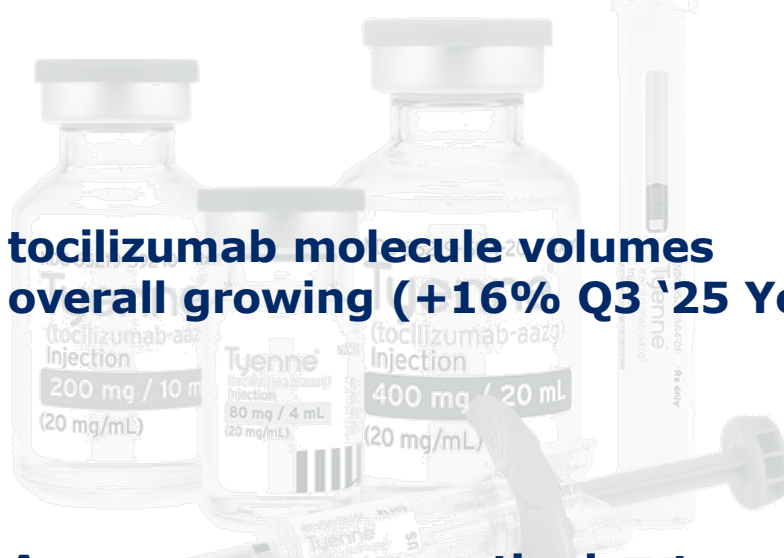
Fastest growing US-launch for pharmacy benefit biosimilar



tocilizumab molecule volumes overall growing (+16% Q3 '25 YoY¹)



Access coverage continuing to expand from parity to exclusive in 2026



Market share¹, in volume %



¹ Source: IQVIA Monthly data



We have a targeted go-to-market-strategy molecule by molecule

Individualized Contracting

Launch learnings and adapting to customer market needs

Customized provider and payer contracting initiatives

Developing partnerships with unique access alternative models

Example: adalimumab-aacf

Grew 74% YoY mostly due to alternative agreements

Access Execution

Market access and contracting strategies built per molecule

Optimizing a brand vs unbranded strategy for lifecycle management

Growing access coverage to bring cost savings to the market

Example: Tylene

Secured formulary placement with payers, now covering >70% of US market and growing

Capability Evolution

Building a hybrid approach in the market for account management

Aligning field and marketing resources to specific initiatives

Flexible capabilities based on molecule, market, and life cycle

Example: Molecule specific

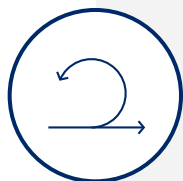
Activate internal teams and resources to support payers/providers to optimize pull through



Track record of agreements to bring volume to market quickly



Proven access agreements
across U.S. regional payers
and provider networks,
distributors



Flexible contracting model
enabling rapid biosimilar
adoption and direct
distribution at scale



**Demonstrated ability to
capture volume early** post-
launch through high-trust,
cost-efficient supply model

Successful innovative agreements

Partner	Scope & impact
	Agreement expanding biosimilar coverage to ~20m members across multiple regional payers
	Our products added to Cost Plus formularies, supporting rapid market share growth
	Exclusive distribution agreement for pharmacy dispensed drugs with access to ~100m lives



Commercial excellence: balanced footprint across geographies



We execute successfully across key regions: deep payer access in EU, gaining traction in US, leader in LATAM, scaling access in other regions



We have a targeted go-to-market strategy molecule by molecule and have shown first-to-market ability with Tyenne, the leading tocilizumab biosimilar



We will continue to de-risk our revenue streams through our commercial network and selected partners with milestone payments

VI

Q&A

VII

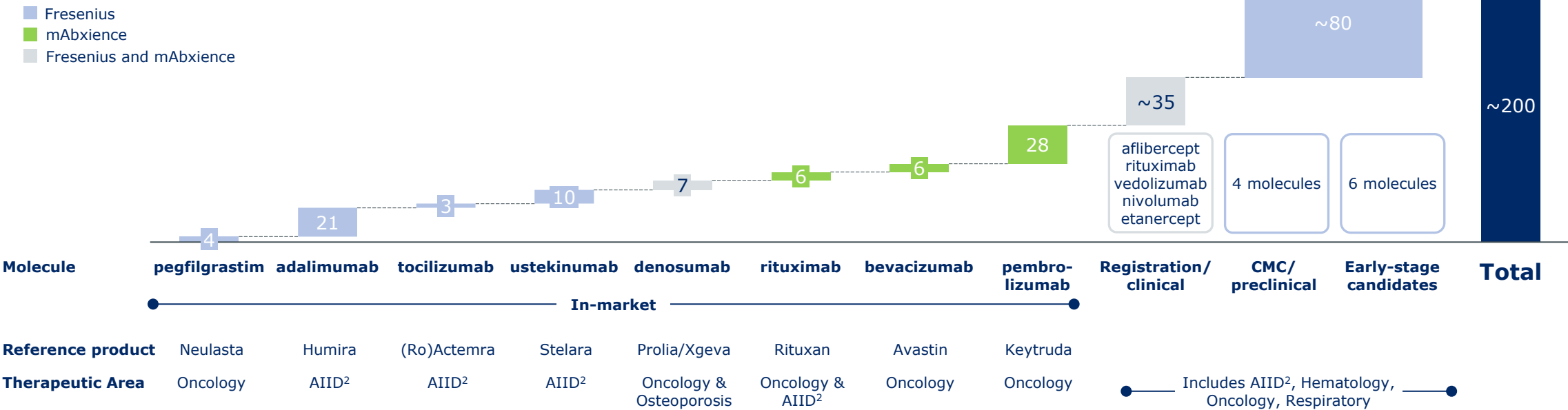
Wrap-up

BACKUP Pipeline & Molecule Onepagers

Our biosimilar portfolio & pipeline

Current biosimilar portfolio & pipeline

Global peak branded sales of originators¹, in €b



> **Attractive and growing biosimilar market** with upcoming near- and mid-term launches

> **Strong position with broad and attractive pipeline**, leveraging end-to-end value chain capabilities

> **Recurring revenues** from milestone payments and CDMO business

¹ Source: Evaluate Pharma | ² Autoimmune & Inflammatory Diseases | ³ Source: IQVIA

Our biosimilar portfolio (in-market)

Limited ●●● Moderate ●●● High ●●●

	Molecule (brand)	Therapeutic Area	Originator peak sales ¹ , €b	Launch			Revenue contribution
				EU	US	RoW	
Fresenius	pegfilgrastim 	Oncology	4	2022	2023		●●●
	adalimumab 	Immunology	21	2019	2023	2021	●●●
	tocilizumab 	Immunology	3	2023	2024	2024	●●●
	ustekinumab 	Immunology	10	2025	2025	2025	●●●
	denosumab 	Osteoporosis, Oncology	7	2025	2025	2025	●●●
	rituximab ²	Oncology	6			2014	●●●
	bevacizumab ³	Oncology	6	2021	2022	2016	●●●
	pembrolizumab	Oncology	28			2024	●●●
	denosumab ⁴	Osteoporosis, Oncology	7	2025	2025	2025	●●●

¹ Source: Evaluate Pharma | ² Referenced as Novex® in LATAM | ³ Referenced as Alymsys®, among other names, in Europe, US and RoW | ⁴ Indicated for osteoporosis and referenced as Izamby® in Europe, Indicated for oncology and referenced as Denbrayce® in Europe

Our biosimilar pipeline

Limited ●●● Moderate ●●●● High ●●●●●

 **Fresenius**

Molecule	Therapeutic Area	Pre-clinical		Clinical		Originator peak sales ¹ , €b	Revenue contribution
aflibercept	Ophthalmology	Early stage	CMC	Ph 1	Ph 3	10	●●●●
rituximab	Oncology	Early stage	CMC	Ph 1		6	●●●●
vedolizumab	Gastroenterology	Early stage	CMC	Ph 1		7	●●●●
Molecule 6	TA's include: Immunology, Hematology, Oncology, Respiratory	Early stage	CMC				
Molecule 7		Early stage	CMC				
Molecule 8		Early stage	CMC				
Molecule 9		Early stage	CMC				
Molecule 10		Early stage				~80	
Molecule 11		Early stage					
Molecule 12		Early stage					
Molecule 13		Early stage					
Molecule 14		Early stage					
Molecule 15		Early stage					
nivolumab	Oncology	Early stage	CMC	Ph 1		10	●●●●
etanercept	Immunology	Early stage	CMC	Ph 1		7	●●●●


From lab to life

¹ Source: Evaluate Pharma

 **Fresenius**

Molecule Overview | adalimumab



Key information

Originator: Humira

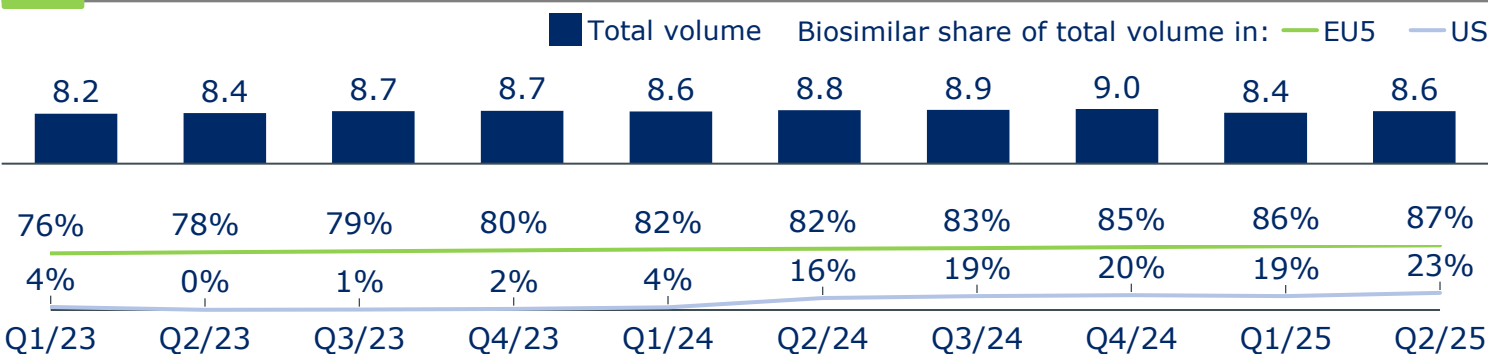
TA: Immunology

Disease: Rheumatoid arthritis, Crohn's disease

Sales contrib.: Moderate

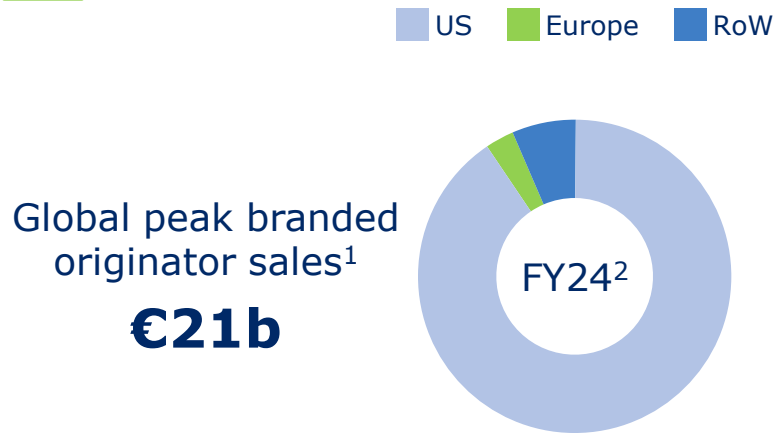
	Launch	# biosimilar products	
		Launched	Pre-Launch
	2019	8	1
	2023	10	

Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

Market size and regional split



- ✓ **Global footprint** in EU, US, LATAM, MENA
- ✓ **Launch of improved TPP ongoing** incl. citrate-free version
- ✓ **In the US, all PBMs kept originator in formulary** in addition to 1-2 biosimilars, resulting in slow initial uptake; acceleration in 2024 after PBM's exclusivity contracting
- ✓ **In the US, primarily reimbursed through Pharmacy Benefit**
- ✓ **Globally, 67% distributed through retail, 33% through hospitals²**

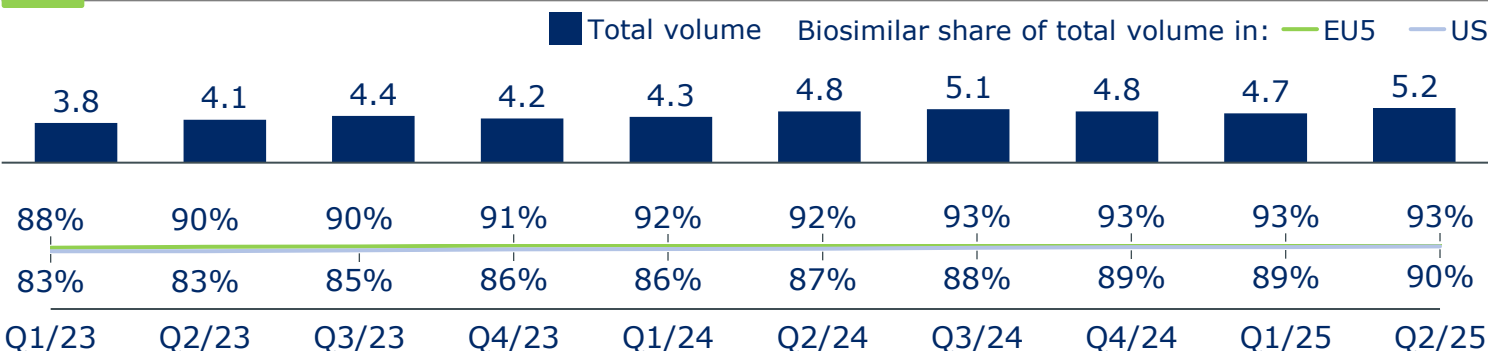
Molecule Overview | bevacizumab

Key information

Originator: Avastin  
TA: Oncology
Disease: Metastatic colorectal cancer
Sales contrib.:    Moderate

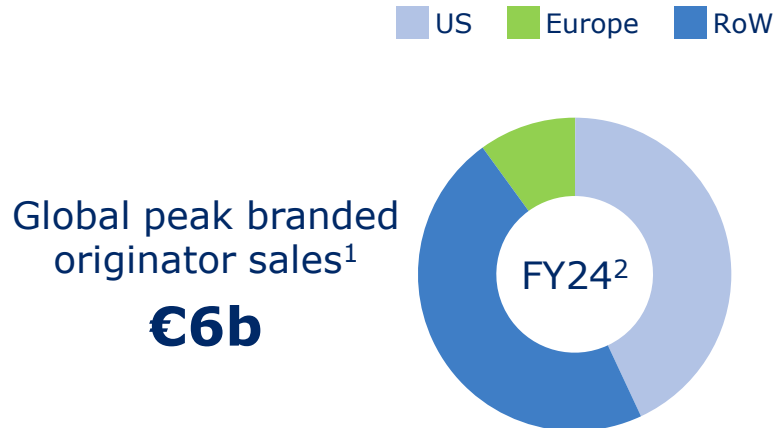
	Launch	# biosimilar products	
		Launched	Pre-Launch
	2021	9	1
	2022	4	1

Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

Market size and regional split



- ✓ **Developed by mAbxience** and marketed globally by several partners
- ✓ **In the US, primarily reimbursed through Medical Benefit**
- ✓ **Globally, 5% distributed through retail, 95% through hospitals²**

Molecule Overview | pegfilgrastim



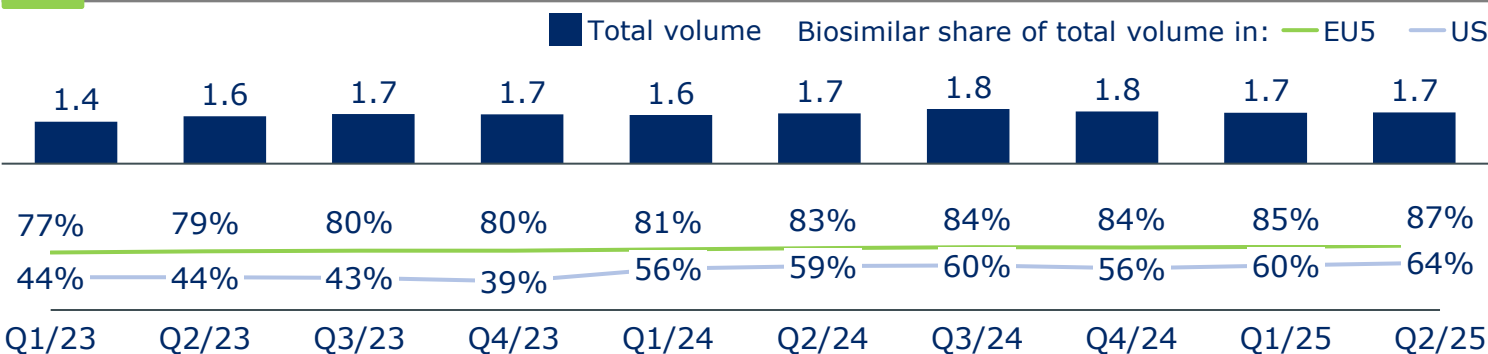
Key information

Originator: Neulasta 
TA: Oncology
Disease: Neutropenia

Sales contrib.:  Limited

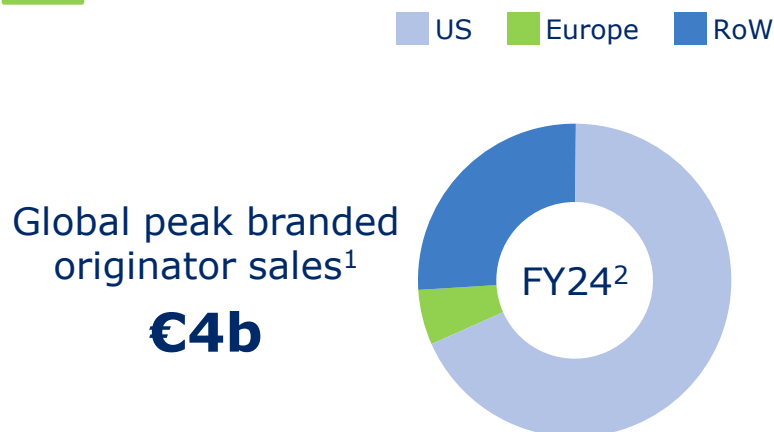
	Launch	# biosimilar products	
		Launched	Pre-Launch
	2022	8	1
	2023	6	1

Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

Market size and regional split



- ✓ In the US, primarily reimbursed through Medical Benefit
- ✓ Globally, 80% distributed through hospital, 20% through retail²

Molecule Overview | tocilizumab



Key information

Originator: Actemra **Genentech**

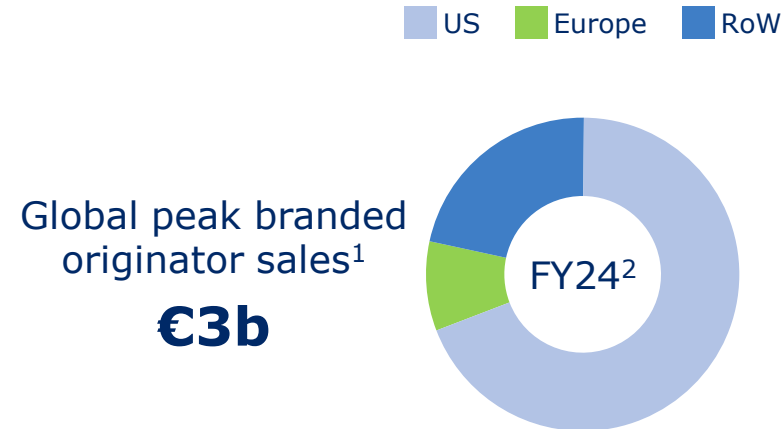
TA: Immunology

Disease: Rheumatoid arthritis

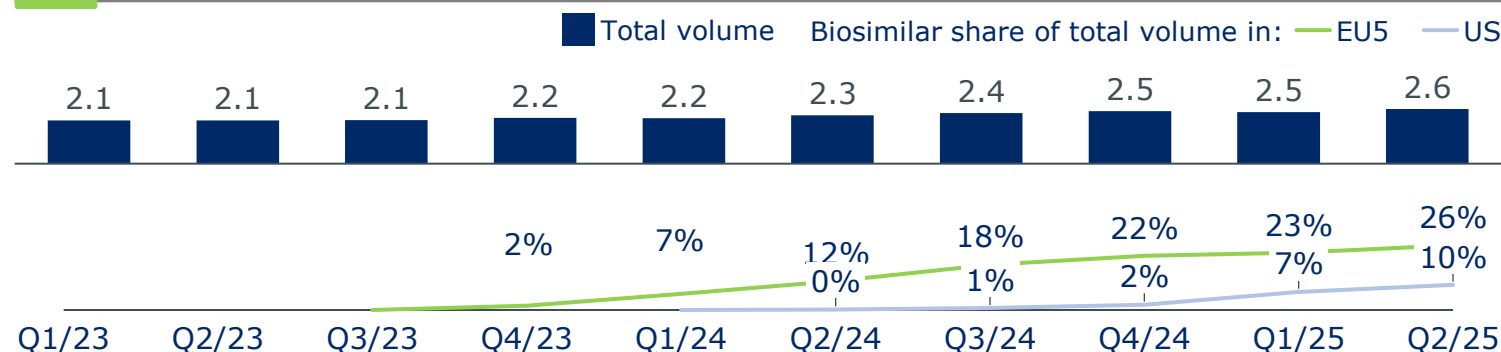
Sales contrib.: ●●● High

	Launch	# biosimilar products	
		Launched	Pre-Launch
	2023	2	2
	2024	3	

Market size and regional split



Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

- ✓ **22 markets launched** – leveraging Autoimmune commercial infrastructure
- ✓ **First-to-market launch**
- ✓ First and only **subcutaneous and intravenous** formulation in the market so far
- ✓ **Advancing with tech transfer** to mAbxience; Garín site received EMA approval
- ✓ **In the US, distribution mixed between pharmacy and medical benefit**
- ✓ **Globally, 51% distributed through hospital, 49% through retail²**

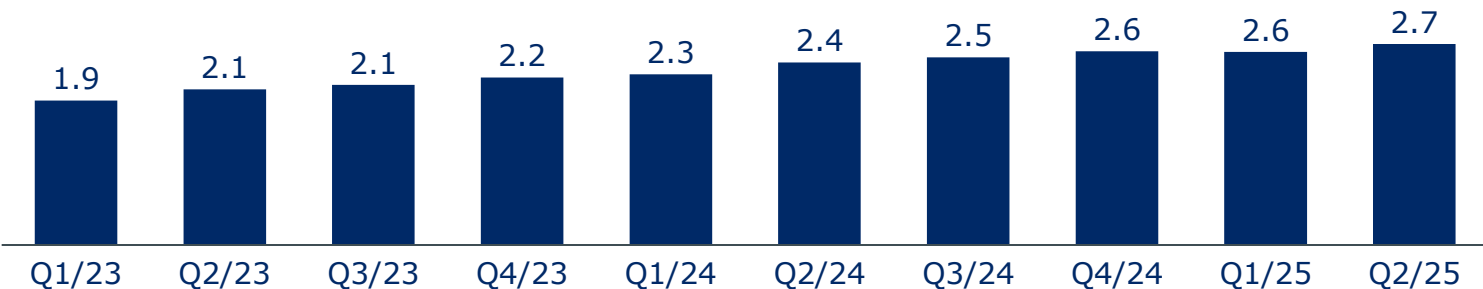
Molecule Overview | pembrolizumab

Key information

Originator: Keytruda 
TA: Oncology
Disease: Unresectable or metastatic melanoma
Sales contrib.:   Moderate

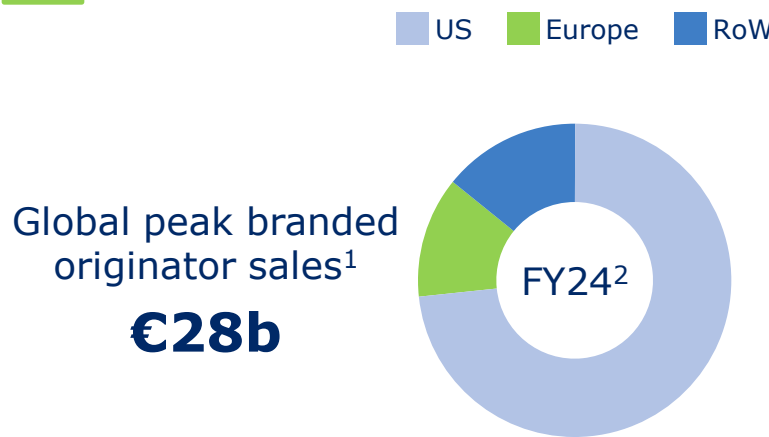
	# biosimilar products	
	Launched	Pre-Launch
		7
		7

Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

Market size and regional split



- ✓ **First-to-market** launch in Argentina and Paraguay in 2025
- ✓ **Developed by mAbxience** and commercialized by Elea ARG and Bioéticos PY
- ✓ **In the US, primarily reimbursed through Medical Benefit**
- ✓ **Globally, 7% distributed through retail, 93% through hospital²**

Molecule Overview | ustekinumab

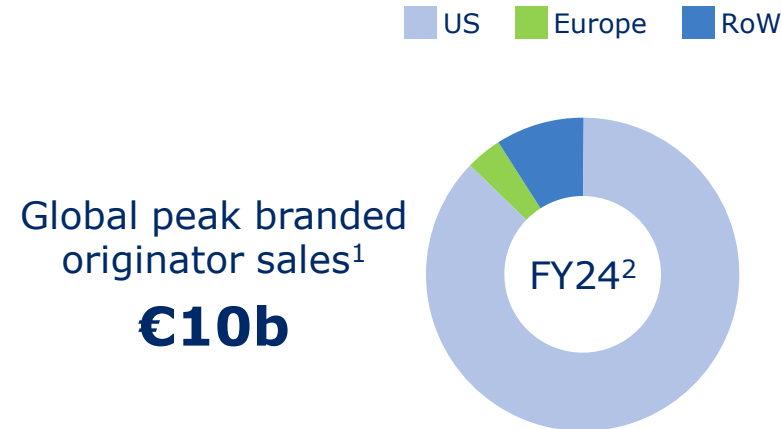


Key information

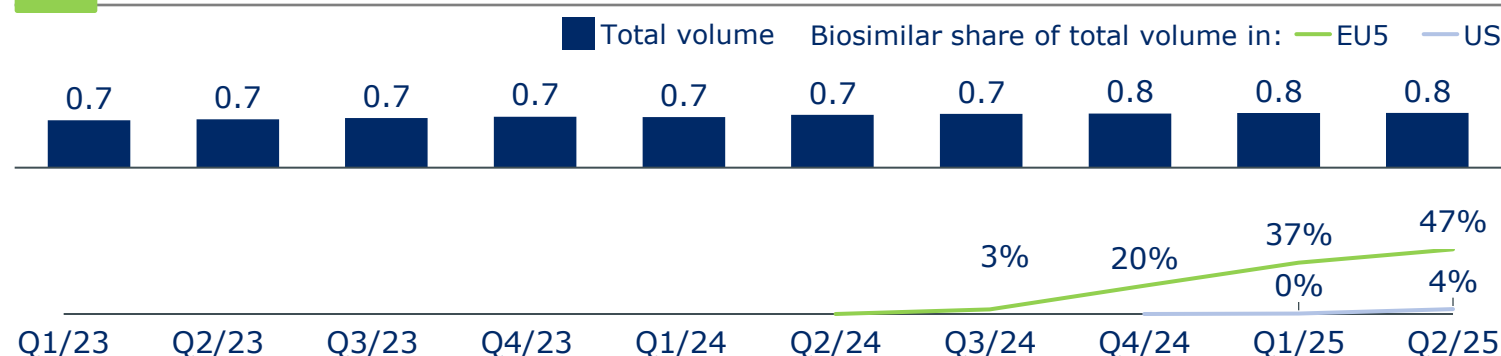
Originator: Stelara 
TA: Immunology
Disease: Plaque psoriasis, Crohn's disease
Sales contrib.:    Limited

	Launch	# biosimilar products	
		Launched	Pre-Launch
	2025	8	1
	2025	8	

Market size and regional split



Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

- ✓ **13 markets launched** – leveraging existing commercial infrastructure
- ✓ **Subcutaneous and intravenous** formulation
- ✓ **U.S. interchangeability** designation
- ✓ **In the US, primarily reimbursed through pharmacy benefit**
- ✓ **Globally, 56% distributed through hospital, 44% through retail²**

Molecule Overview | denosumab



Key information

Originator: Prolia / Xgeva

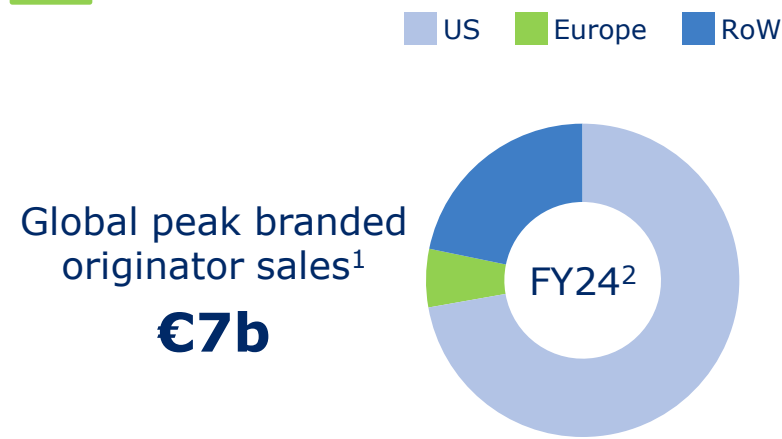
TA: Osteoporosis, Oncology

Disease: Osteoporosis, cancer-related bone loss

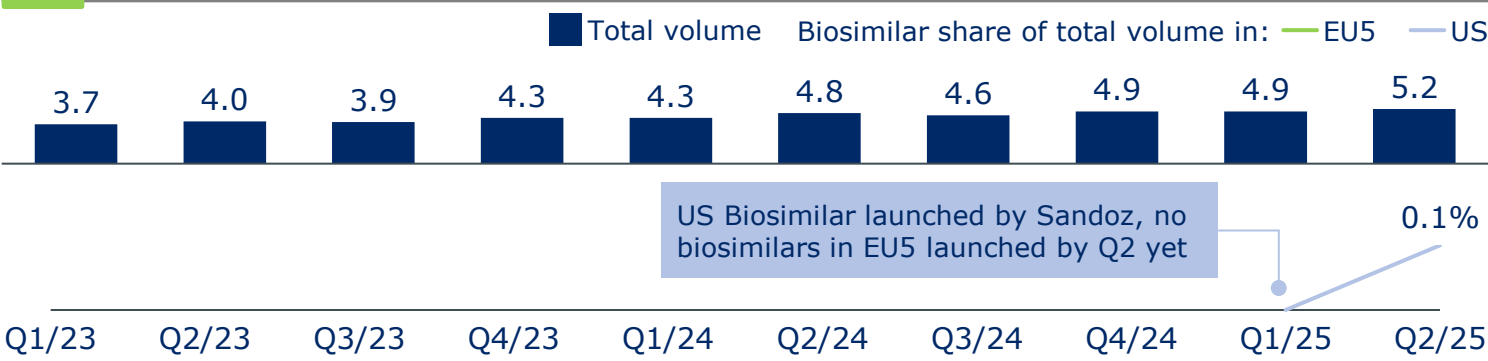
Sales contrib.: Moderate

	Launch	# biosimilar products	
		Launched	Pre-Launch
	2025	5	8
	2025	3	8

Market size and regional split



Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

- ✓ **First wave to market** with large TPP offering
- ✓ **Differentiated vs competitors** through Oncology PFS
- ✓ **For US launch**, CMS³ issued a permanent, product-specific billing code
- ✓ **In the US, reimbursement balanced** between retail and hospital
- ✓ **Globally, 59% distributed through retail**, 41% through hospital²

Molecule Overview | aflibercept

Key information

Originator: Eylea **REGENERON** 

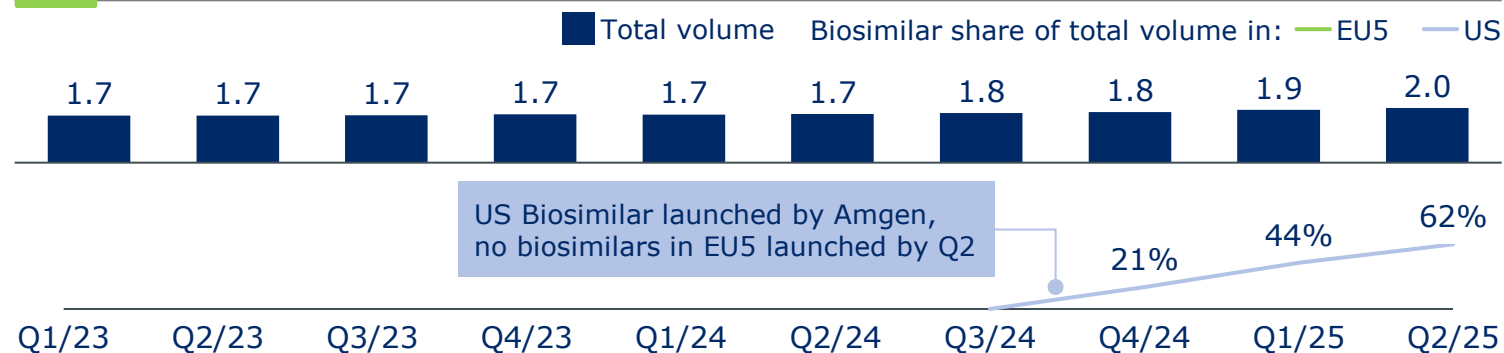
TA: Ophthalmology

Disease: Neovascular age-related macular degeneration

Sales contrib.:  Moderate

	# biosimilar products	
	Launched	Pre-Launch
		8
	1	7

Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

Market size and regional split

Global peak branded originator sales¹

€10b

Aflibercept not well covered in IQVIA, e.g., underestimation of US market size

✓ **In-licensed from SamChunDang Pharm** with exclusive commercialization for US and several LATAM countries

✓ **In the US, primarily reimbursed through Medical Benefit**

✓ **Globally, 44% distributed through retail, 56% through hospitals²**

Molecule Overview | vedolizumab

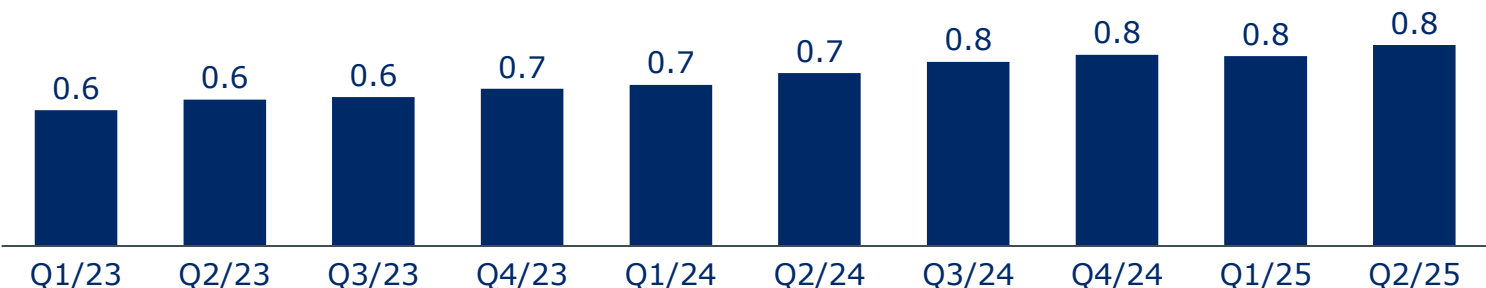
Key information

Originator: Entyvio 
TA: Gastroenterology
Disease: Ulcerative colitis, Crohn's disease

Sales contrib.: ●●● High

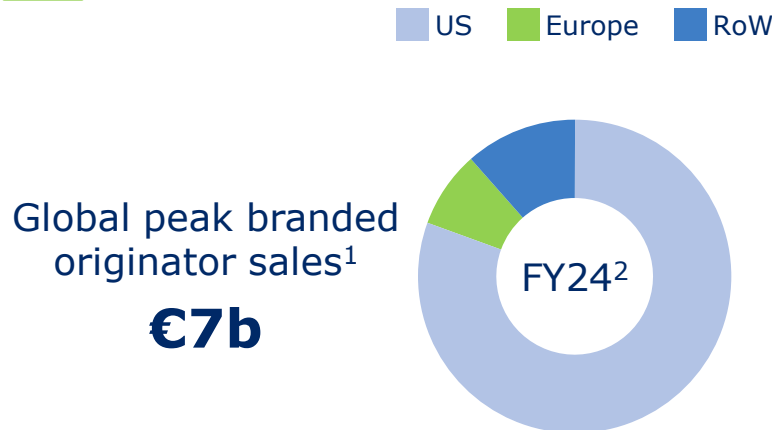
	# biosimilar products	
	Launched	Pre-Launch
		3
		4

Volume, in m standard units²



¹ Source: Evaluate Pharma | ² Source: IQVIA

Market size and regional split



✓ **Global in-licensing agreement announced with Polpharma Biologics**

✓ **In the US, reimbursement mixed between pharmacy and medical Benefit**

✓ **Globally, 44% distributed through retail, 56% through hospital²**

Glossary

Glossary (1/4)

Term	Explanation
Aseptic FP capabilities	Ability to manufacture finished pharmaceutical products in sterile environments to prevent microbial contamination.
ASP	Average Selling Price; the average price at which a drug is sold to purchasers, often used in reimbursement calculations
ASP+ / ASP+ reimbursement model	A U.S. Medicare payment model that reimburses providers at the Average Selling Price plus an additional percentage to cover handling costs.
Auto-injector	A prefilled, self-administered device designed to deliver a fixed dose of medication via subcutaneous or intramuscular injection.
Bioreactor	A vessel or system that provides a controlled environment for growing cells or microorganisms to produce biologic substances.
Cell line	A population of cells derived from a single cell with uniform genetic makeup used for consistent biologic production.
cGMP / GMP	Current Good Manufacturing Practice; regulatory standards ensuring products are consistently produced and controlled for quality and safety.
Citrate-free formulation	A drug formulation that removes citrate buffer to reduce injection-site pain and improve patient comfort.
Clone	A cell or group of cells derived from a single parent cell with identical genetic material used in biologic production.
CMC	Chemistry, Manufacturing, and Controls; the technical section of drug development describing product composition and quality assurance processes.
CMS	Centers for Medicare & Medicaid Services, a U.S. federal agency that administers the major healthcare programs and regulates, among others, healthcare reimbursement, pricing, and data reporting.
Cost of Production (COP)	The total expense incurred in manufacturing a drug, including raw materials, labor, utilities, equipment, and overhead, often used to benchmark competitiveness and profitability in biomanufacturing
Digital Twins	Virtual digital replicas of physical processes or systems used to simulate, monitor, and optimize biomanufacturing performance.
Drug Product (DP)	The finished dosage form of a pharmaceutical containing the active ingredient(s) and excipients, ready for patient use.

Glossary (2/4)

Term	Explanation
Drug Substance (DS)	The active pharmaceutical ingredient that provides the intended therapeutic effect in a medicine.
ELN	Electronic Laboratory Notebook; a digital system for recording, storing, and sharing laboratory data and experiments.
EMA	European Medicines Agency; the regulatory authority responsible for the scientific evaluation and supervision of medicines in the EU.
FDA	U.S. Food and Drug Administration; the regulatory agency overseeing safety, efficacy, and quality of drugs, biologics, and medical devices.
Finished Product (FP)	The final packaged pharmaceutical product released for commercial distribution.
Formulary	A list of medications approved for use and reimbursement by a particular health plan, hospital, or payer.
IDN (Integrated Delivery Network)	A network of healthcare providers and facilities under shared management aiming to coordinate patient care and reduce costs.
In-licensing	Acquiring the rights to develop or commercialize a product or technology originally developed by another organization.
Interchangeability	A regulatory designation allowing a biosimilar to be substituted for its reference product without prescriber intervention.
Intravenous (IV)	Administration of a drug directly into a vein for rapid systemic delivery.
Latex-free	Indicates materials and packaging are free of natural rubber latex to prevent allergic reactions.
Lead clone	The selected cell line or genetic variant demonstrating optimal yield and product quality for scale-up manufacturing.
LIMS	Laboratory Information Management System; software for managing laboratory samples, workflows, and analytical data.
LoE (Loss of Exclusivity)	The point when a branded drug's patent protection expires, allowing generic or biosimilar competition. LoE-1 refers to the year before.

Glossary (3/4)

Term	Explanation
Medical benefit	Insurance coverage for drugs administered by healthcare professionals, typically billed under the medical plan.
Medicare Advantage	U.S. Medicare Part C plans offered by private insurers providing combined hospital and medical coverage, sometimes including drugs.
Out-licensing	Granting rights to another company to market a product owned by the licensor.
Parallelization of clinical trials	Conducting multiple clinical activities concurrently to shorten overall development timelines.
Payer	An organization, such as an insurance company or government agency, that finances or reimburses healthcare costs.
PBM	Pharmacy Benefit Manager; intermediaries that negotiate drug prices and manage prescription benefits on behalf of payers.
Pharmacy benefit	Coverage for outpatient prescription drugs dispensed through retail or specialty pharmacies.
Phase 1	The first stage of clinical trials assessing safety, tolerability, and pharmacokinetics in healthy volunteers or patients.
Phase 3 waiver	Regulatory allowance to approve a biosimilar without a traditional Phase 3 trial based on robust analytical and PK/PD similarity data.
PK / PD	Pharmacokinetics and Pharmacodynamics; the study of how the body affects a drug and how the drug affects the body.
PMDA	Pharmaceuticals and Medical Devices Agency; Japan's authority for evaluating and approving drugs and medical devices.
Pre-Filled-Syringe (PFS)	A single-use syringe preloaded with a sterile drug product, ready for administration.
Process validation	Documented proof that a manufacturing process consistently produces product meeting predetermined quality standards.
Q-Code	A billing code used in the U.S. Medicare system for drugs administered under the medical benefit, often for biologics.

Glossary (4/4)

Term	Explanation
Regulatory track record	A company's historical performance and reliability in obtaining and maintaining regulatory approvals.
Scale-up	Increasing production from laboratory or pilot to commercial manufacturing while maintaining product quality.
Single-use technology	Disposable bioprocessing equipment designed to prevent cross-contamination and reduce cleaning needs.
Small-scale similarity	Early comparative testing of biosimilar and reference product on limited scale to confirm analytical equivalence.
Subcutaneous (SC)	Administration of a drug into the fatty tissue beneath the skin for slower absorption.
Target Product Profile (TPP)	A strategic summary outlining desired characteristics of a drug product to guide development and regulatory alignment.
Tech transfer	The process of transferring manufacturing or analytical knowledge and processes between development and production sites.
Tender	A formal procurement process where suppliers submit bids to provide medicines at negotiated prices to public buyers.
Therapeutic Area (TA)	A specific category of diseases or medical conditions targeted by related treatments, such as oncology or immunology.
Titer	The concentration of product (e.g., antibody) produced by cells in a bioreactor, typically measured in grams per liter.
Transfection	Introduction of foreign genetic material into cells to enable production of a desired protein or biologic.
Vial	A small glass or plastic container used to store and administer injectable drug products.
Yield	The amount of product obtained from a bioprocess relative to input materials, reflecting production efficiency.

