



**JUBILANT
PHARMOVA**

July 29, 2025

BSE Limited,
Floor 25, P. J. Towers
Dalal Street, Fort
Mumbai - 400 001

Scrip Code: 530019

National Stock Exchange of India Limited,
Exchange Plaza, Bandra-Kurla Complex,
Bandra (E),
Mumbai - 400051

Symbol: JUBLPHARMA

Sub: Press Release alongwith Earnings Presentation on the financials and operational performance of the Company for the quarter ended June 30, 2025

Ref: Regulation 30 of SEBI (Listing Obligations and Disclosure Requirements) Regulations, 2015 ("Listing Regulations")

Dear Sirs,

Pursuant to Provisions of Regulation 30 of the Listing Regulations, please find enclosed herewith the Press Release, Presentation and FAQs on the financials and performance of the Company for the quarter ended June 30, 2025.

The above mentioned documents will be simultaneously posted on the Company's website at www.jubilantpharmova.com.

You are requested to kindly take the same on record.

Thanking you,

Yours faithfully,
For Jubilant Pharmova Limited

Naresh Kapoor
Company Secretary

Encl: as above

A Jubilant Bhartia Company

OUR VALUES



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www.jubilantpharmova.com

PRESS RELEASE

Noida, Jul 29, 2025

JUBILANT PHARMOVA – Q1' FY26 RESULTS

On track towards Vision 2030

Solid growth momentum along with EBITDA & PAT margin expansion

Strong customer traction in CDMO Sterile Injectable business

Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue from operations	1,732	1,929	1,901	10%
Total Income	1,746	1,941	1,913	10%
EBITDA	266	357	302	14%
<i>EBITDA Margin (%)</i>	<i>15.2%</i>	<i>18.4%</i>	<i>15.8%</i>	<i>60 bps</i>
Normalised PAT¹	69	139	103	48%
Normalised PAT Margin	4.0%	7.1%	5.4%	140 bps

1. Normalised PAT is after adjusting for exceptional items and tax.

In Q1'FY25, Reported PAT at Rs.482 Cr. was higher due to one-time net exceptional income of Rs. 396 Cr.

The Board of Jubilant Pharmova Limited met today to approve financial result for the quarter ended June 30, 2025.

Commenting on the Company's performance in Q1'FY26, **Mr. Shyam S Bhartia, Chairman Jubilant Pharmova Limited and Mr. Hari S Bhartia, Co-Chairman & Non-Executive Director, Jubilant Pharmova Limited** said, "We are pleased to announce revenue of Rs. 1,901 Cr. in Q1'FY26, growth of 10% over last year, same quarter. It is heartening to note is that we delivered solid revenue growth across all of our business units and we expect this growth momentum to grow stronger as we move forward. EBITDA grew by 14% YoY to Rs. 302 Cr. Ebitda margins expanded by 60 basis points on the back of improved operating performance across CRDMO and Generics. Normalised PAT grew by 48% to Rs. 103 Cr. on the back of improved operating performance and reduced finance cost. Reported PAT in Q1'FY25 at Rs. 482 Cr. was higher because of one time net exceptional income of Rs. 396 Cr. As we are consciously investing in Radiopharma, CDMO Sterile Injectables and CRDMO business to secure future growth, Net Debt / EBITDA increased marginally from 1.1x in Mar'25 to 1.2x in Jun'25 on the back of increased capex intensity."

During Q1'FY26, we saw continued growth momentum from Ruby-Fill® and PET radiopharmacies. In the Allergy Immunotherapy, we witnessed increase in demand from the US. In the CDMO Sterile Injectables, strong customer traction for Line 3 in Spokane continues. In the CRDMO business, we integrated the new R&D facility in France and are now investing in business development initiatives. In the Generics business, we are foreseeing growth & profitability improvement. Lastly, in our Proprietary Novel drugs business, we continue to progress in dosing patients in JBI-802 and JBI-778 clinical trials."

Q1'FY26 Financial Highlights

In Q1'FY26, Revenue grew by 10% on a YoY basis to Rs. 1,901 Cr. on the back of growth in revenue across all business units. EBITDA grew by 14% on a YoY basis to Rs. 302 Cr. due to improved performance in CRDMO and Generics. Q1'FY26 normalised PAT increased by 48% on a YoY basis to Rs. 103 Cr. on the back of improved operating performance and reduced finance cost. Reported PAT in Q1'FY25 at Rs. 482 Cr. was higher because of one time exceptional income of Rs. 396 Cr.

Segmental Business Performance

Radiopharma - *Leading Radiopharmaceutical manufacturer & 2nd largest Radiopharmacy network in the US*

Radiopharmaceuticals Q1'FY26 revenue grew by 3% to Rs. 271 Cr. and EBITDA remained stable YoY at Rs. 126 Cr. The business continues to maintain a strong position in the high margin SPECT imaging product portfolio. On the PET side, The Ruby-Fill® installations are increasing. We are on track to introduce multiple new products in the PET and SPECT imaging from FY27 to FY29. The dosing for Phase 2 clinical trial for MIBG is complete and we are preparing data package to be submitted to FDA by H2'FY26.

Radiopharmacy Q1'FY26 revenue grew by 5% YoY to Rs. 598 Cr. EBITDA margins for Q1'FY26 stands at 2%. EBITDA margins remained weak due to increased competitive intensity in the SPECT business. In H2'FY25, two of our PET radiopharmacies have started distributing PYLARIFY®, which is an industry leading prostate cancer diagnostic imaging agent. We continue to see increase in revenue from PET radiopharmacies.

The proposed investment of US\$ 50 million in PET radiopharmacy network is underway. This investment will take the overall PET radiopharmacy network to Nine (9) sites, thereby solidly positioning Jubilant Pharmova's radiopharmacy network as the second largest in the US and shall drive the future business growth.

Allergy Immunotherapy - *No. 2 in the US Sub-Cutaneous allergy immunotherapy market*

As the sole supplier of Venom in the US, the business is expanding the overall market by increasing customer awareness. In the US Allergenic extracts, the business is working to increase revenues. The business is also working to increase penetration in the outside US markets.

In Q1'FY26, revenues grew by 8% to Rs. 181 Cr. on the back of growth in revenues from US market. EBITDA remained stable YoY at Rs. 63 Cr. EBITDA margin for Q1'FY26 stands at 35%. We anticipate outside US sales to gradually improve.

CDMO Sterile Injectables – *Leading contract manufacturer in North America, serving top global innovators*

Q1'FY26 revenue grew by 14% to Rs. 370 Cr. due to increase in sales volume. EBITDA grew by 9% to Rs. 62 Cr. EBITDA margins are lower QoQ due to annual maintenance shutdown at Spokane facility. The capacity expansion program in Spokane, Washington, USA is on track. Media fills had been successfully completed on Line 3 and the technology transfer programs are underway. The large innovator pharma companies are now looking to create an alternate manufacturing site in the US as a risk management strategy to mitigate any potential tariff's imposed by the US govt. In light of the same, we are starting to see excellent traction in Requests for Proposals (RFPs) for Line 3 including from Big Pharma. We expect to finalise these within FY26. The commercial production on line 3 is expected to start in FY26. We also expect to reach peak utilisation for Line 3 in three years post start of commercial production vs four years, expected earlier. The Montreal facility continued operations after successful implementation of corrective and preventive actions.



CRDMO – Indian leader for integrated drug discovery & formidable API player

In Q1'FY26, the Drug Discovery business revenue grew by 42% to Rs. 161 Cr. EBITDA grew by 46% to Rs. 32 Cr. Revenue continue to increase due to increase in revenue from large Pharma customers. We have integrated new R&D facility in France and are now investing in business development. EBITDA margins lower QoQ due to change in project mix and investment in business development. Overall, the medium term outlook continues to be positive on the back of the increase in large pharma clients and the addition in new capabilities.

The API business revenue grew by 9% to Rs. 141 Cr in Q1'FY26. EBITDA grew by 36% to Rs. 22 Cr. EBITDA margins improved by 310 basis points due to profitable product mix.

We have proposed sale and transfer of API Business to Jubilant Biosys Limited, a wholly owned subsidiary of the Company. This transaction will result in housing of the drug discovery business and CDMO API business in a single business entity. This combined platform will improve the operational efficiency in the business and lead to superior brand recall of “Jubilant Biosys Limited” as provider of end-to-end CRDMO services by the large pharmaceutical & Biotech customers. The transaction will also help to improve asset utilisation of API business by improving the revenue mix towards Custom manufacturing & CDMO.

Generics – Building a growing, profitable & agile business model

In Q1'FY26, the Generics business revenue grew by 7% to Rs. 166 Cr. EBITDA for the period stands at Rs. 12 Cr. Revenue increase is primarily driven by Non US markets. EBITDA margins improved by 1,400 basis points YoY due to focus on profitable products.

We plan to launch 6 to 8 products per annum in our US and non-US international markets. In line with our plan, we are ramping up exports to the US markets in a meaningful and gradual manner. We have also started supply of products from our Contract manufacturing partners to the US market.

Proprietary Novel Drugs – Innovative biopharmaceutical company developing breakthrough therapies

The global clinical trials for our lead programs, Phase II trial for JBI -802 for Essential Thrombocythemia (ET) and other Myeloproliferative Neoplasms (MPN) and Phase I trial for JBI -778 for non-small cell lung cancer (NSCLC) and high grade Glioma are actively enrolling patients and progressing in line with our expectations.

About Jubilant Pharmova Limited

Jubilant Pharmova Limited (formerly Jubilant Life Sciences Limited) is a company with a global presence that is involved in Radiopharma, Allergy Immunotherapy, CDMO Sterile Injectables, Contract Research Development and Manufacturing Organisation (CRDMO), Generics and Proprietary Novel Drugs businesses. In the Radiopharma business, the Company is involved in the manufacturing and supply of Radiopharmaceuticals with a network of 45 radiopharmacies in the US. The Company's Allergy Immunotherapy business is involved in the manufacturing and supply of allergic extracts and venom products in the US and in some other markets such as Canada, Europe and Australia. Jubilant Pharmova Limited through its CDMO Sterile Injectables business offers manufacturing services including sterile fill and finish injectables (both liquid and lyophilization), full-service ophthalmic offer (liquids, ointments & creams) and ampoules. The CRDMO business of the Company includes the Drug Discovery Services business that provides contract research and development services through two world-class research centers in Bengaluru and Noida in India and one in France. The CDMO-API business that is involved in the manufacturing of Active Pharmaceutical Ingredients. Jubilant Therapeutics is involved in the Proprietary Novel Drugs business and is an innovative biopharmaceutical company developing breakthrough therapies in the area of oncology and auto-immune



disorders. The Company operates multiple manufacturing facilities that cater to all the regulated markets including USA, Europe and other geographies. Jubilant Pharmova Limited has a team of around 5,500 multicultural people across the globe. The Company is well recognised as a 'Partner of Choice' by leading pharmaceuticals companies globally.

For more information, please contact:

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Disclaimer

Statements in this document relating to future status, events, or circumstances, including but not limited to statements about plans and objectives, the progress and results of research and development, potential product characteristics and uses, product sales potential and target dates for product launch are forward-looking statements based on estimates and the anticipated effects of future events on current and developing circumstances. Such statements are subject to numerous risks and uncertainties and are not necessarily predictive of future results. Actual results may differ materially from those anticipated in the forward-looking statements. Jubilant Pharmova may, from time to time, make additional written and oral forward looking statements, including statements contained in the company's filings with the regulatory bodies and its reports to shareholders. The company assumes no obligation to update forward-looking statements to reflect actual results, changed assumptions or other factors.



JUBILANT
PHARMOVA

Earnings Presentation Q1'FY26

Disclaimer



Statements in this document relating to future status, events, or circumstances, including but not limited to statements about plans and objectives, the progress and results of research and development, potential product characteristics and uses, product sales potential and target dates for product launch are forward looking statements based on estimates and the anticipated effects of future events on current and developing circumstances. Such statements are subject to numerous risks and uncertainties and are not necessarily predictive of future results. Actual results may differ materially from those anticipated in the forward-looking statements. Jubilant Pharmova may, from time to time, make additional written and oral forward looking statements, including statements contained in the company's filings with the regulatory bodies and our reports to shareholders. The company assumes no obligation to update forward-looking statements to reflect actual results, changed assumptions or other factors.

Jubilant Bhartia Group has created value across multiple sectors



Strong presence in diverse sectors

- Pharmaceuticals
- Life Science Ingredients
- Performance Polymers
- Food Service (QSR)
- Contract Research & Development Services
- Therapeutics
- Auto Dealerships
- Oil and Gas services



Global presence through investments

- India
- USA
- Canada
- Europe
- Singapore
- Australia
- Africa
- China
- Sri Lanka, Bangladesh



Employer of Top Talent

43,000 people across the globe with ~2,200 in North America

Jubilant Pharmova, a diversified pharmaceutical company



Radiopharma

Leading manufacturer

of Radiopharmaceuticals
in North America

2nd largest radiopharmacy network in the US



Allergy Immunotherapy

2nd largest player

in the US Allergenic extract market
Sole supplier of Venom
Immunotherapy in the US



CDMO Sterile Injectables

Leading contract manufacturer

in North America
Serves top global innovator pharma
companies



CRDMO

Integrated drug discovery

and development service provider
Formidable API player
in multiple therapeutic areas



Generics

Over 50 countries served

including regulated markets
Broad therapeutic areas :
CVS, CNS, GI and MS



Proprietary Novel Drugs

Two drug programs

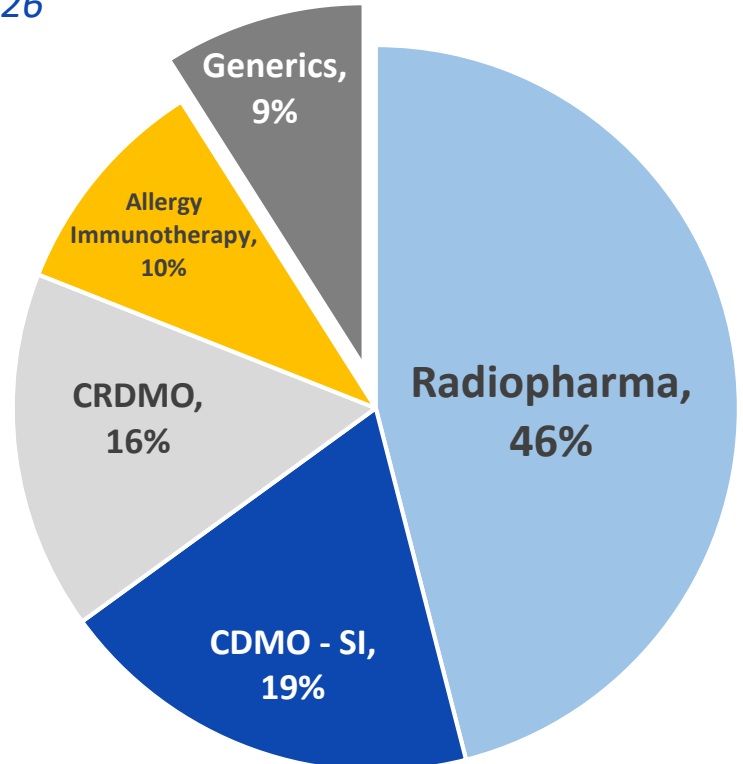
in clinical trials
Developing high potential precision
medicines in Oncology

A global leader with a
strong team of 5,500
people

Focus on specialty products & services and Dollar revenues

Business wise Revenue Split

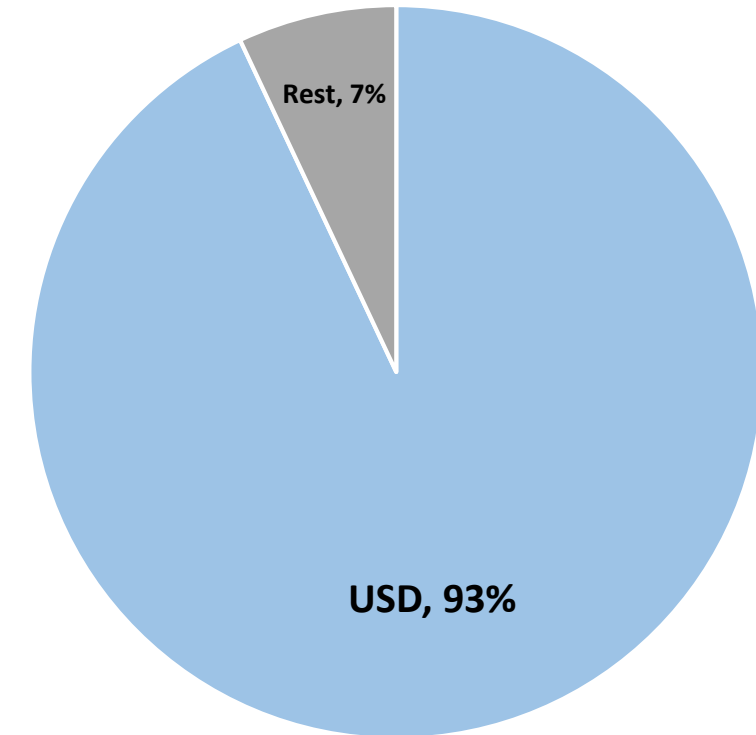
Q1'FY26



Specialty Products (Radiopharma, Allergy Immunotherapy) and Specialty Services (CDMO & CRDMO) contribute majority of revenues

Currency wise Revenue Split

Q1'FY26

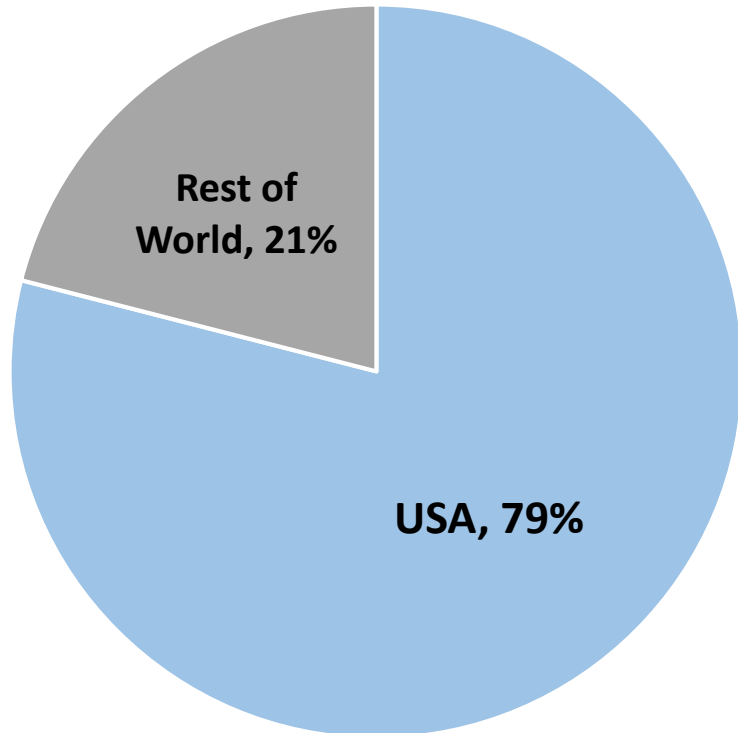


Majority revenues are USD denominated

Minimal risk from US Tariffs

Geography wise Revenue Split

Q1'FY26



US market constitutes
majority of revenues

Origin of Goods & Services sold in the US

Q1'FY26



Goods from Canada (Radiopharmaceuticals) exempted from tariffs under US- Canada – Mexico trade agreement

* Goods and Services from Canada 17% : Goods 16%, Services 1%

* Goods and Services from India 15% : Goods 7%, Services 8%

State-of-the-art manufacturing and research facilities enable our growth



NORTH AMERICA



INDIA & EUROPE



G. Noida, Uttar Pradesh - Drug discovery



Bengaluru, Karnataka - Drug discovery



France - Drug discovery

6
Manufacturing facilities

3
Research facilities

45
Radiopharmacies

Vision 2030: We aspire to double our revenues by FY30 and we are on the right track



From
FY24

To
FY30

*Actual
Trailing 12 Months*



2X Revenue

Rs. 6,703 Cr.

Rs. 13,500 Cr.

Rs. 7,404 Cr.

25% EBITDA Margin

~ 15 %

23% to 25%

17%

Zero Net Debt

Rs. 2,457 Cr.

Zero

**Rs. 1,535 Cr.
End of Q1'FY26**

High Teens RoCE

High Single digit

High Teens

**12%*
Q1'FY26 Annualised**

• (EBIT before exceptional items) / Average ((Equity + Gross Debt) less (CWIP adjusted for grant))

These are our growth drivers to achieve Vision 2030



Business	Growth Drivers
Radiopharma	Leadership in Ruby-Fill® Launch New PET, SPECT and Therapeutic products (MIBG) Invest in 6 high margin PET Radiopharmacies in US
Allergy immunotherapy	Strengthen competitive position and develop new products
CDMO - Sterile Injectables	Double capacity in Spokane, US
CRDMO	Add large pharma customers Grow CDMO and custom manufacturing in API
Generics	Launch new products in the US and Grow profitable Non-US international business



Radiopharma

Radiopharmaceuticals



SPECT
Imaging

Low Energy
gamma rays
detected by SPECT cameras



PET
Imaging

High Energy
positrons
detected by a PET scanner



Radiopharmaceutical
Therapeutics

**Systemically or Locally
Delivered**
radiation using pharmaceuticals

Isotopes - Tc99m

Isotopes - Rb82, F18, Ga68

Isotopes – I131, Lu177, Ac225

Key Products

MAA, DTPA, Sulfur Colloid,
Mertiatide

Ruby-Fill®, Pylarify, Illuccix,
Neuraceq, FDG

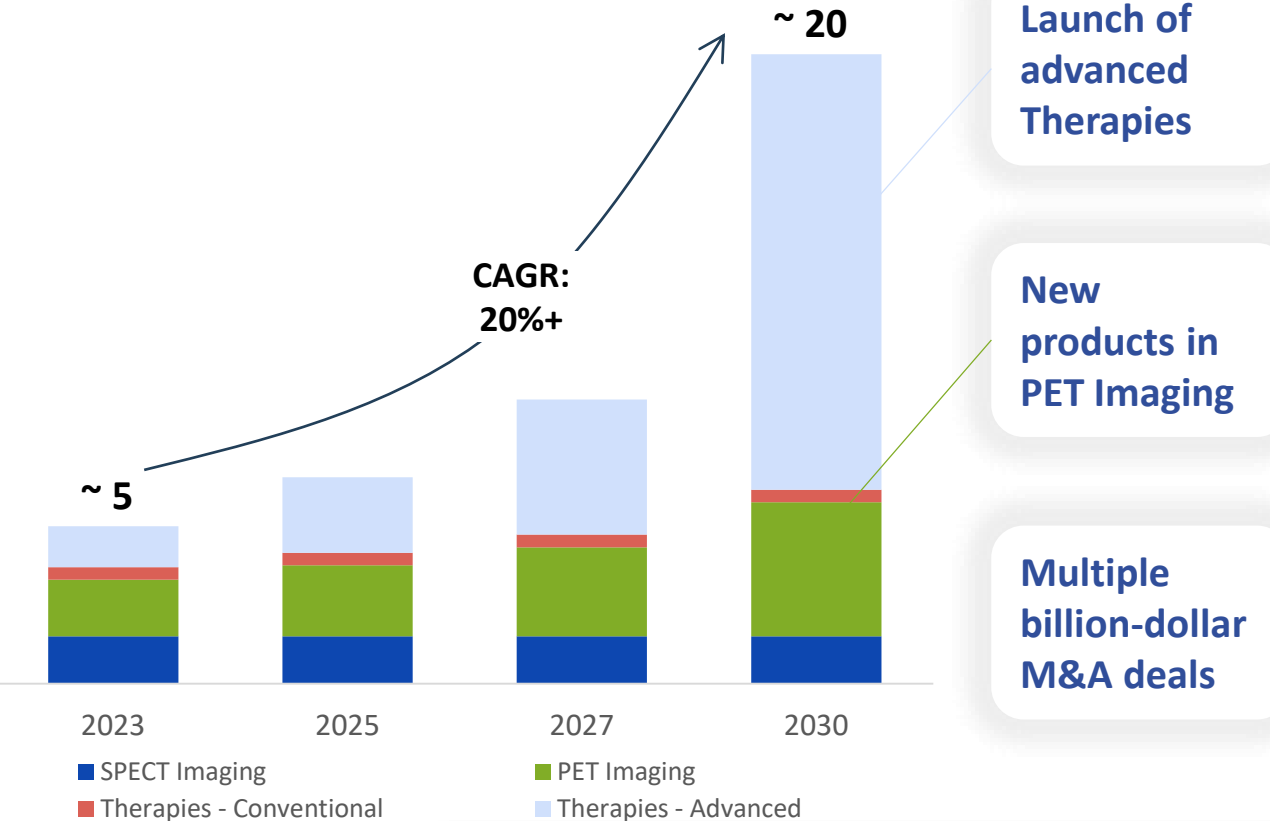
HICON® Sodium Iodide
I 131, Pluvicto, Lutathera

Radiopharmaceuticals have a growing role in treatment of life-threatening diseases
e.g. Cancer

US Radiopharmaceutical market is growing at 20% CAGR

US Radiopharmaceutical Market

USD Bn.



Growth Drivers & Trends

- PSMA Therapeutic, Pluvicto for Prostate Cancer ~USD 2.0 Bn.
- PSMA Diagnostics for Prostate Cancer ~ USD 1.5 Bn.
- Broad range of applicability e.g. Alzheimer's
- Special reimbursement for diagnostic products (FIND Act)
- Novartis and Mariana Oncology (USD 1 Bn.)
- AstraZeneca and Fusion (USD 2.4 Bn.)
- Lilly and Point Biopharma (USD 1.4 Bn.)
- BMS and Rayzebio (USD 3.6 Bn.)

PET imaging & advance therapies are driving the market growth

Consolidated Market with high Entry Barriers



Managing time sensitive logistics

Radioactive isotope decays exponentially. The half life could be few hours to few days. Goal is to deliver high activity doses

Stringent manufacturing & regulatory environment

Adherence with **extensive license framework**. Stringent manufacturing set up required to handle isotopes

Forward integration with radiopharmacies

Forward integration with radiopharmacies **helps to gain market share**

Innovative new product development

High capex requirement, long developmental cycle and **complex isotope handling requirements** for novel product development.

We are a leading Radiopharmaceuticals manufacturer in North America



	Organ	Key Indication	Product
PET Dx	Cardiac	Coronary Artery disease	Ruby - Fill®
	Breast	Lymph nodes detection	Sulfur Colloid
	Cardiac	Cardiac blood pool imaging	Tc99m-Glueceptate
		Coronary Artery Disease	Tc99m-Sestamibi
SPECT Dx	Gastrointestinal	Intra-abdominal Infection	Tc99m-Exametazime
	Lung	Pulmonary Embolism	Tc99m-DTPA
		Pulmonary Perfusion	Tc99m-MAA
	Muscoskeletal	Altered osteogenesis	Tc99m-MDP
	Renal	Renal failure	Tc99m-Mertiatide
	Thyroid	Localising thyroid malignancies	I-131
Therapeutics	Thyroid	Hyperthyroidism, Thyroid Cancer	I-131 HICON®

- Diversified across diagnostics & therapeutics
- Current TAM at USD 400 Mn.
- Strong R&D and supply chain
- In-house API manufacturing

Market leadership in select products

Draximage® MAA



Draximage® DTPA



Ruby-Fill®



HICON® Sodium Iodine I 131
Solution USP



MAA is used in the **perfusion** phase of a ventilation/perfusion (V/Q) scan to diagnose **pulmonary embolism**. JDI is leading player in the US market

DTPA is used to assess **pulmonary ventilation function** in association with MAA to perform a Ventilation/perfusion (V/Q) scan. JDI is leading player in the US market

It is used for Cardiac PET scan, to evaluate regional myocardial perfusion in adults with suspected or existing coronary artery disease. **JDI is the innovative leader in the US market**

HICON® is a radioactive therapeutic agent indicated for the treatment of hyperthyroidism and selected cases of carcinoma of the thyroid. **JDI has no direct competition in the US market**

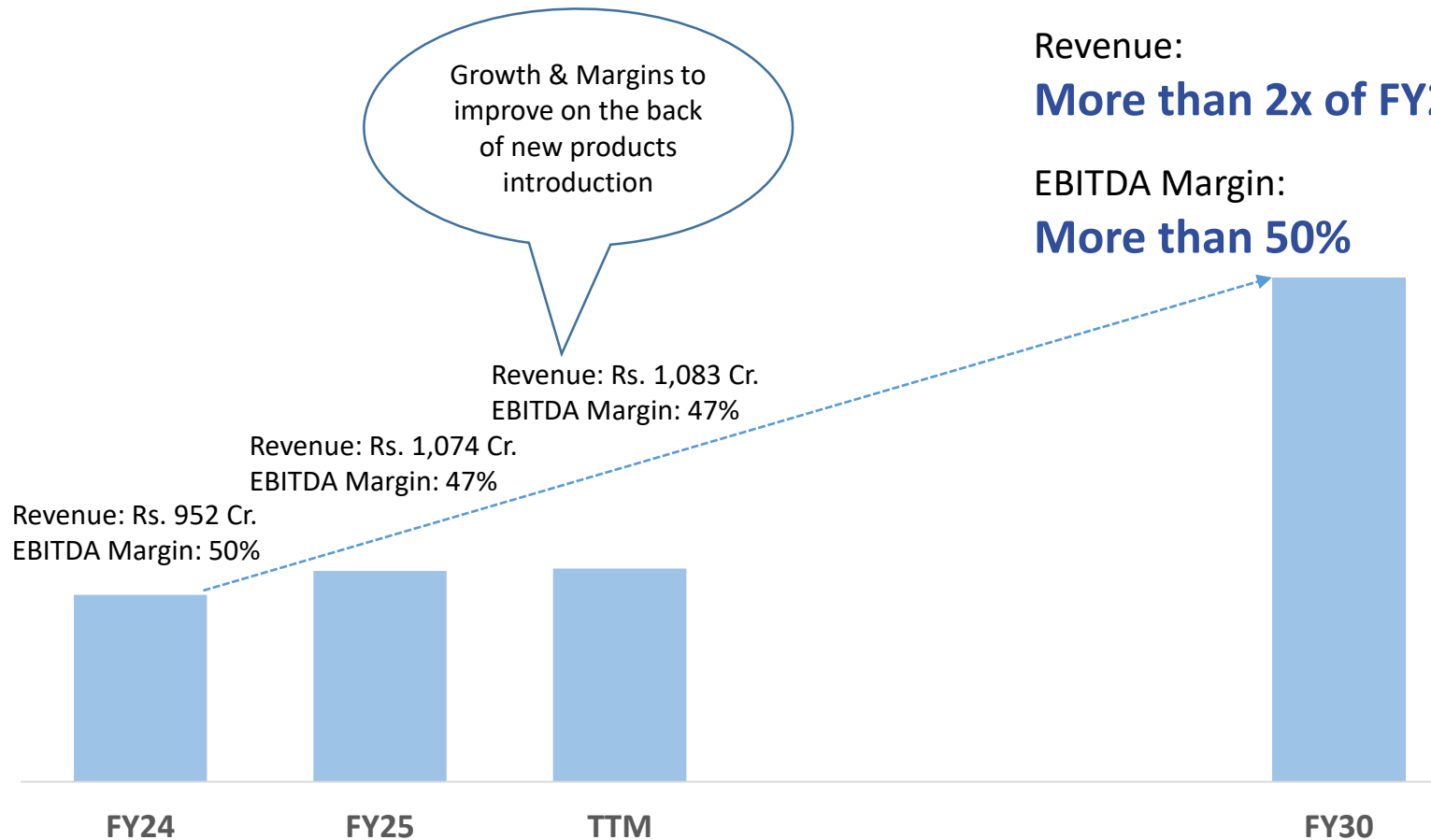
Radiopharmaceuticals Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	262	296	271	3%
EBITDA	126	136	126	0%
<i>EBITDA Margin (%)</i>	48%	46%	46%	(150) bps

- Q1'FY26 revenue grew on back of growth in Ruby-Fill ® and Sulfur Colloid
- Q1'FY26 EBITDA stable YoY

Radiopharmaceuticals Vision 2030: To more than double the revenues



Growth drivers:

- Ruby-Fill®
- New PET & SPECT products
- MIBG

To become leader in cardiac PET Imaging through Ruby-Fill®



Ruby-Fill ® Rubidium 82 generator and Elusion System

Growth drivers:

- Ruby-Fill®
- New PET & SPECT products
- MIBG



Competitive advantage

- Longer life per generator (7 weeks vs 6 weeks for peer)
- Better image quality and consistency
- Constant Activity

Current Position

- Market Size ~ USD 180 Mn. and growing at 12%
- Market share ~ 25% and growing

Product Innovation

- Value engineering to lower cost & improve margin
- AI enabled 3D cardiac blood flow quantification

Launch new PET and SPECT imaging products with a TAM of USD 550 Mn



Developing new products in SPECT Imaging to maintain leadership & in PET Imaging for growth

Growth drivers:

- Ruby-Fill®
- New PET & SPECT products
- MIBG



Timeline	Incremental TAM USD Mn.	Potential Peak Annual Sales - USD Mn.	No. of launches
FY27	50	20	2
FY28	250	60	3
FY29	250	40	4
Total	550	120	9

Launch MIBG by FY27

Growth drivers:

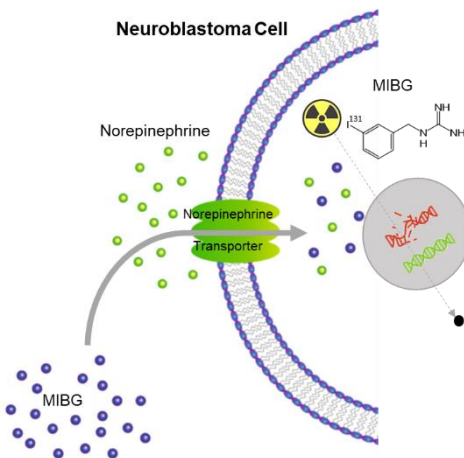
- Ruby-Fill®
- New PET & SPECT products
- MIBG

HICON® Sodium Iodide I 131 - Commercialised



- Iodine I 131, HICON® is standard care for patients
- Used for diagnosis and treatment of Thyroid cancer
- Used in imaging & treatment for pediatric cancer - Neuroblastoma
- Relapsed / Refractory patients have limited treatment options

MIBG - Undergoing Clinical trials



- Potential peak sales USD 70 - 100 Mn.
- Data package to FDA by H2'FY26



Radiopharmacy

Radiopharmacies are critical in generating value

SPECT Radiopharmacy



PET Radiopharmacy

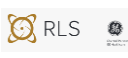


Growth Drivers & Trends

- Consolidated market in the US. Large M&A transactions in Radiopharmacies
- Increasing demand for novel PET products driving PET radiopharmacies growth
- Stringent USP 825 regulations to drive increase in therapeutics dispensing through Pharmacy
- Emerging radioisotopes landscape such as Rb-Sr, Ga-68, Cu-64, Lu-177, Ac-225

Consolidated market with high Entry Barriers

Consolidated Market

	# of radio pharmacies in the US	SPECT pharmacies	PET pharmacies	# of hospitals served in the US
 CardinalHealth™	160+	✓	✓	~ 4,100
 JUBILANT RADIOPHARMA	45	✓	✓	~ 1,800
 SIEMENS Healthineers PETNET Solutions	41		✓	~ 700
 RLS	31	✓		~ 900
 PharmaLogic Take The Lead	42	✓	✓	~ 200
 SOFIE	14		✓	~ 200

Barriers to Entry

1

Stringent Regulations

Each treatment site is required to obtain a license from Nuclear Regulatory Commission and comply with additional state, local, and hospital regulations for transportation and usage

2

Intricate Supply Chain

A robust supply chain is required given short product half-lives and strong customer preference for just-in-time ordering, compared to large bulk orders

3

Complex Care Coordination

Requires awareness, education, and collaboration across multiple hospital departments

4

Skilled Manpower Requirement

Authorized nuclear pharmacists require at least 4,000 hours of training or experience in nuclear pharmacy practice along with rigorous examinations

The 2nd largest radiopharmacy network in the US



45

Radiopharmacies
with ~20%
volume market
share



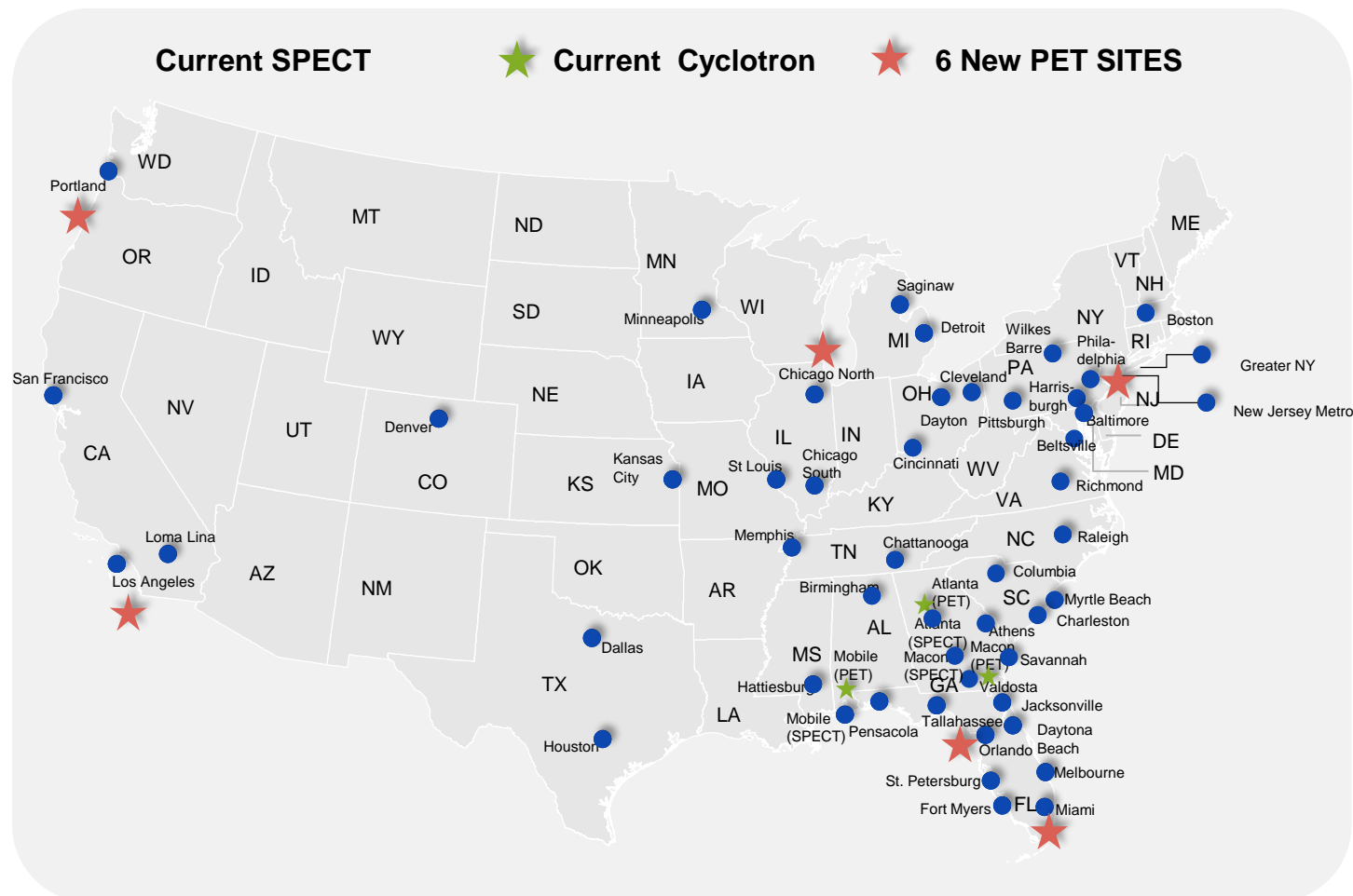
1,800
hospitals
catered



6 customized
doses delivered
every
minute



99%+
on-time deliveries,
Use of AI for route
optimization



USP<825>

JDR network is USP 825
compliant



Business moat

Unique combination of
SPECT manufacturing &
radiopharmacy network



6

Planning new sites in
PET network



Therapeutics

distribution is preferred
from radiopharmacies

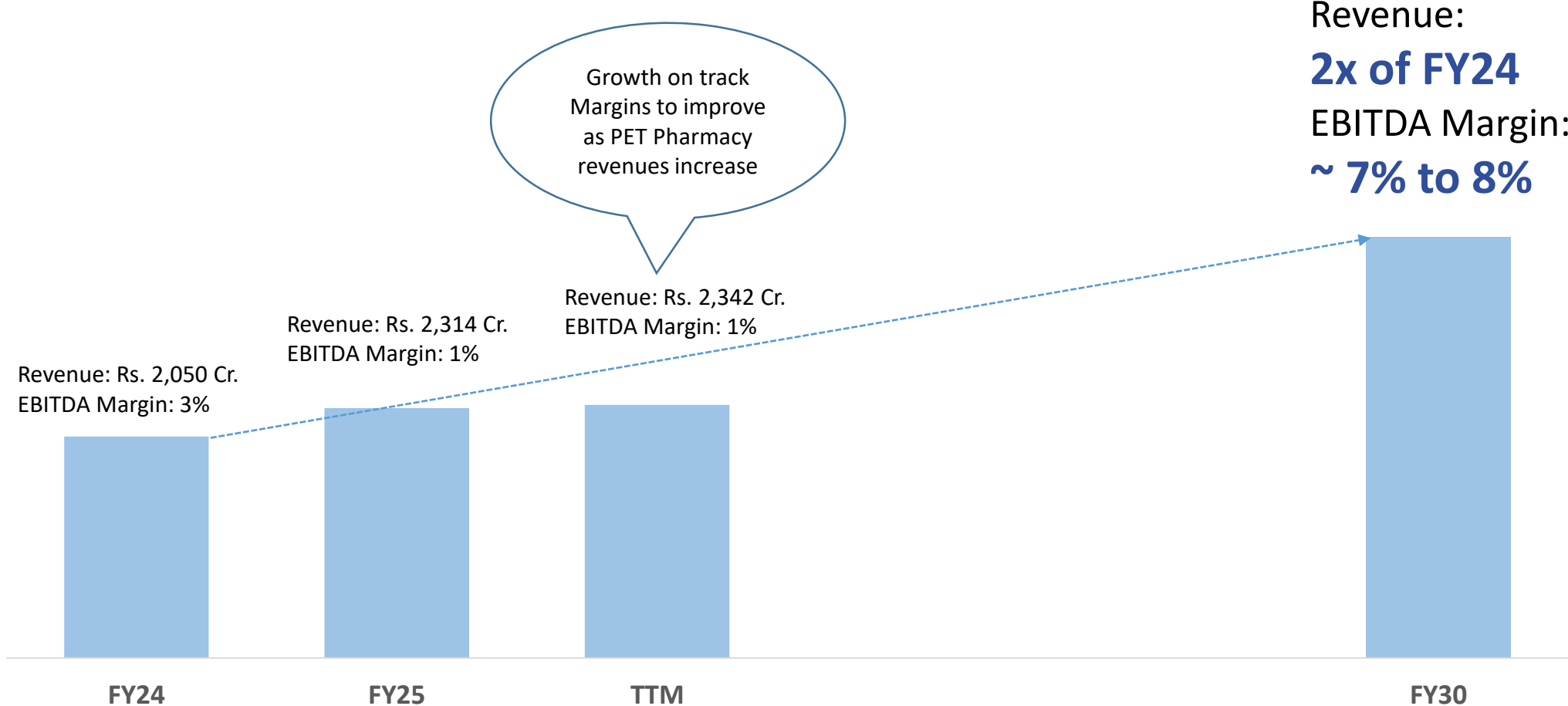
Radiopharmacy Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	570	600	598	5%
EBITDA	13	6	10	(22%)
<i>EBITDA Margin (%)</i>	2%	1%	2%	(60) bps

- Q1'FY26 revenue grew YoY on the back of increase in volume from new PET products
- Q1'FY26 EBITDA lower YoY due to increase in competitive intensity in SPECT radiopharmacies

Radiopharmacy Vision 2030: Double the revenues, expand margins by adding 6 PET Radiopharmacies



Expanding PET Radiopharmacy network from current 3 sites to 9 sites



Growth driver:

- PET expansion



- Strengthened network to enable long term contracts** with PET radiopharmaceutical manufacturers
- Fully operational by FY28.** Funding through internal accruals and long-term credit
- Expect Asset turnover of 1.0x and RoCE 20% +** on the USD 50 Mn. investment

Continue to increase in PET radiopharmacy revenues from the current 3 sites



Allergy Immunotherapy

Allergy immunotherapy is the sole way to fundamentally reduce allergen hypersensitivity

- 20% + global population have allergies e.g. Asthma and Allergic Rhinitis
- Allergy Immunotherapy requires repeated shots of allergic antigens to develop immunity

Allergies



Testing



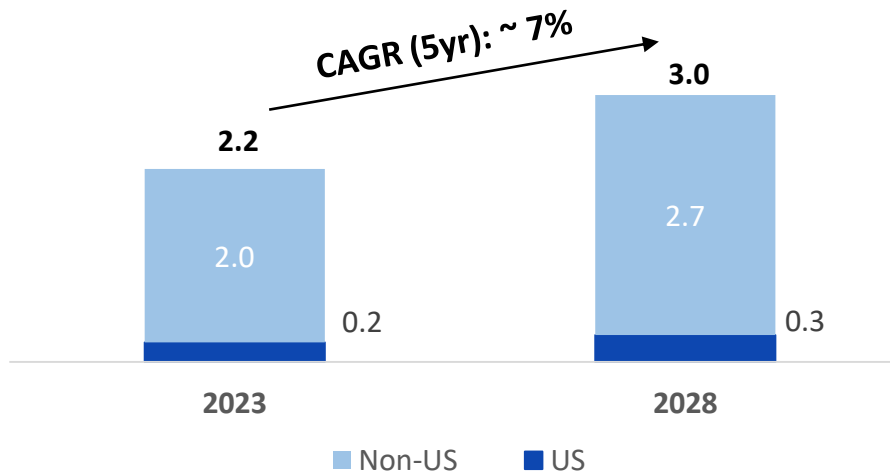
Treatment



Global Allergy Immunotherapy market is expected to grow by ~ 7%



Global Allergy Immunotherapy Market
USD Bn.



Growth Drivers and Trends

- Concentrated US market with 3 players
- Complex supply chain from sourcing to processing
- Grandfathered approvals, new product needs BLA
- Market increasing in Sub-Lingual delivery
- Challenging reimbursement landscape

2nd largest player in the US Sub-Cutaneous Allergy Immunotherapy market



- 100-year-old 'HollisterStier' brand
- Sole Supplier of Venom extracts in the US
- 200+ allergenic & 6 venom extracts
- Onshore US FDA approved manufacturing
- Dedicated sales force in the US
- 2,000+ Allergists / ENTs as customers

Venom Extracts



Venom extracts for Honey Bee and other insects

Allergenic Extracts



Allergenic extracts for Dog, Cat, Mite, Tree, Pollen etc.

Skin Testing Devices



Multiple skin testing systems

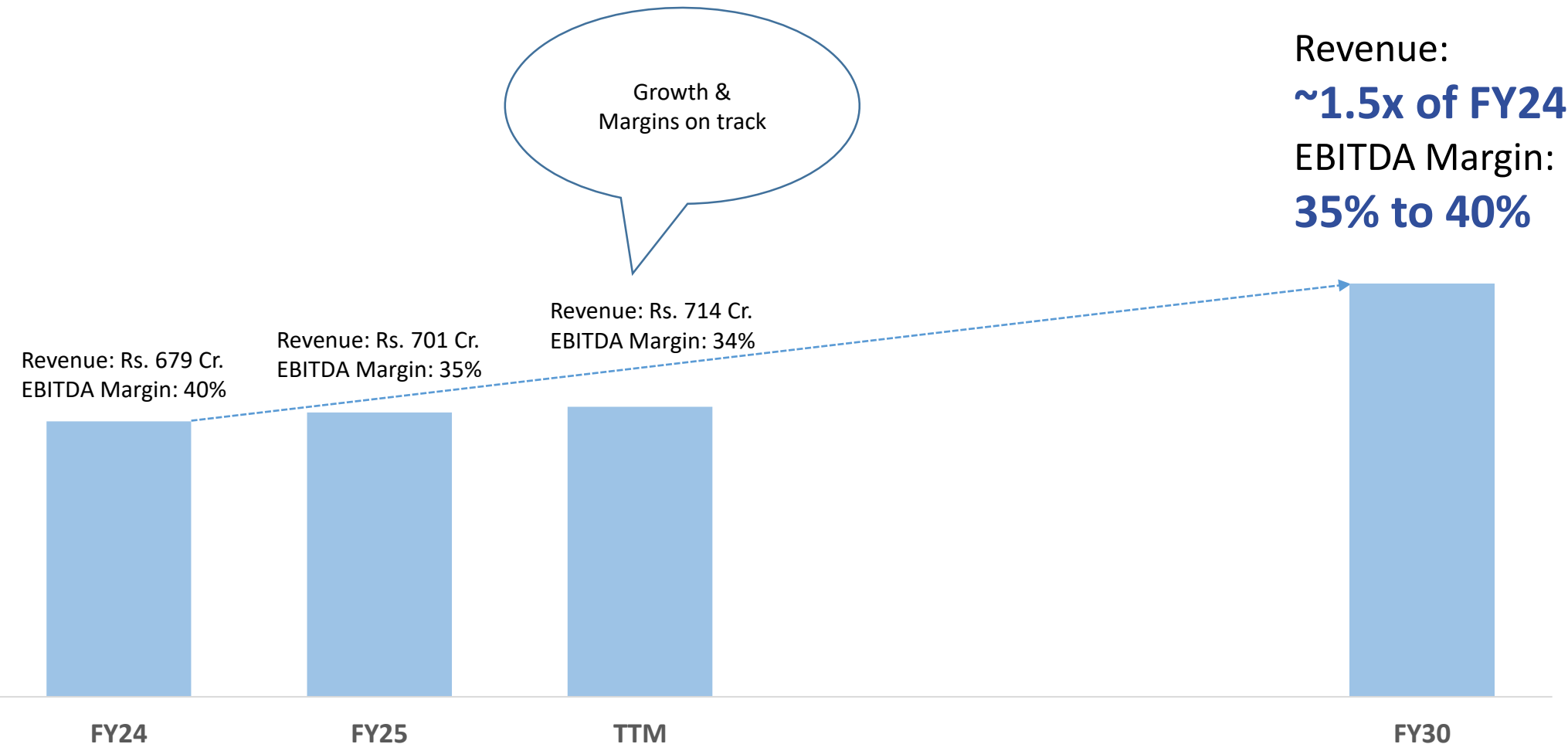
Allergy Immunotherapy Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	168	192	181	8%
EBITDA	63	88	63	(1%)
EBITDA Margin (%)	38%	46%	35%	(300) bps

- Q1'FY26 revenue grew on the back of revenue growth in the US market
- Q1'FY26 EBITDA stable YoY. EBITDA margins in the normalized range

Allergy Immunotherapy Vision 2030: Solidify position as a scientific leader



Allergy Immunotherapy

Growth Drivers



Strengthen competitive position in US

- Retain and grow **Venom customers** & patient base
- Increase **US revenue** in **Allergenic extracts** through targeted marketing

Grow outside US business

- Increase **outside US Venom sales** through strategic partnerships in European markets

Increase investment in R&D

- Develop new products & technologies
- Build treatment innovation through partnerships and alliances





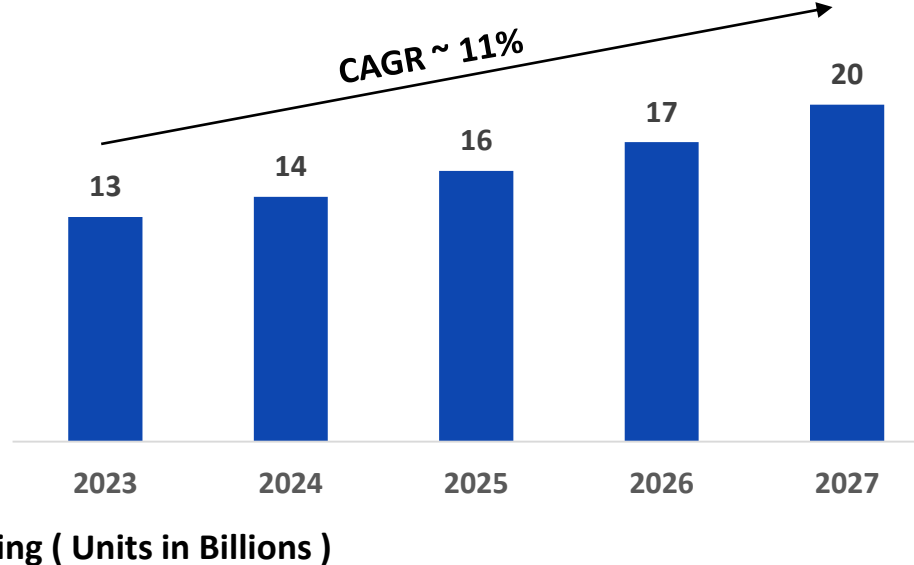
CDMO - Sterile Injectables

CDMO - Sterile Injectables is seeing demand supply gap widening



Global CDMO-SI Market Size

USD Bn



Vial filling (Units in Billions)

Year	2023	2024	2025	2026	2027
Demand	4.9	5.2	5.7	6.2	6.8
Supply	5.5	5.8	6.1	6.1	6.1

Demand supply gap of 700 Mn. vials in 2027, to be further widened by industry consolidation

Growth Drivers & Trends

- Innovator Pharma companies, for their US requirement, are planning to shift the manufacturing from Europe to US, as a risk mitigation measure due to impending Tariffs by the US Govt.
- Consolidation in supply due to large acquisitions - Catalent Inc. by Novo Holding
- Increasing number of drugs in Biologics pipeline and Loss of exclusivity
- Reduction in offshoring by innovators due to regulatory and supply chain advantages

Market with high Entry Barriers



- **Majority of commercial contracts are typically long duration** (typically 3 years or more with auto renewal)
- **Greenfield expansion is considerably difficult** due to high up-front capex required with ongoing opex to support initial product commercialization
- **Innovator companies prefer onshore North American manufacturers** with a good quality track record in light of continuing supply challenges
- **Attractive niches & Technology** (e.g., Isolator Technology, Multi Dose Preservative Free ophthalmic drops, etc.) have emerged, driven by requirements of differentiated technologies, higher quality standards, people capabilities and capital investment
- **High switching costs for customers** due to significant tech transfer time (18-24 months), other challenges, e.g., quality
- **Stringent regulatory requirements (FDA) for sterile manufacturing**, with ever evolving landscape making difficult for new entrants

We are a leading North American CDMO player with unique capabilities and strong customer relationships



- **5 of the top 20** pharma companies as customers
- **25+** customers across the world with multiple products having patent protection and limited competition
- **5+ years** average relationship time with Top 10 Customers
- **90%+ repeat customer** business
- **24 months** of switching timelines for customers
- **Full suite of services** including sterile fill and finish (Liquid & Freeze dried), Ophthalmic (Liquids, Ointments and creams) and Biologics
- **10+ years of US FDA compliant status** at flagship site in Spokane

The business is engaged in Fill and Finish for Sterile Injectables, where a sterile drug is transferred from a filling needle into a sterile vial and then a stopper is applied, except in cases, where the drug requires sterile lyophilization.

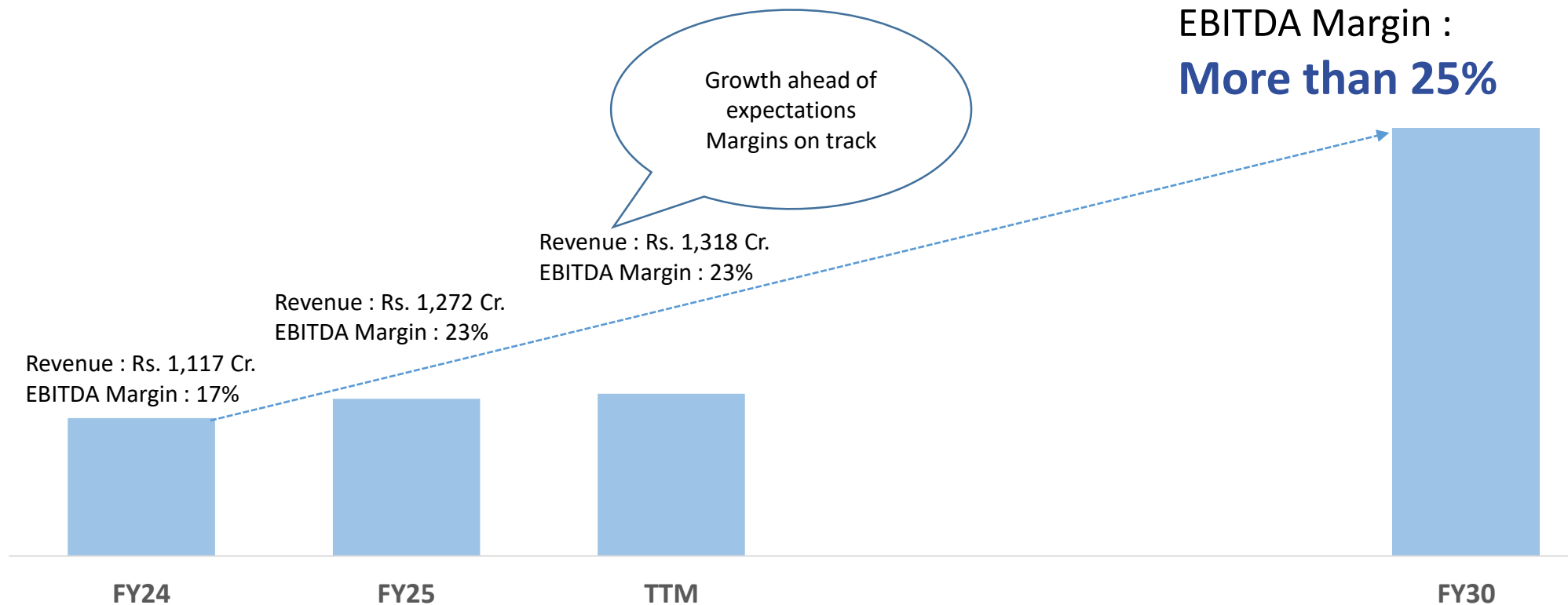
CDMO Sterile Injectables Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	324	340	370	14%
EBITDA	57	95	62	9%
<i>EBITDA Margin (%)</i>	<i>18%</i>	<i>28%</i>	<i>17%</i>	<i>(90) bps</i>

- Q1'FY26 revenue grew YoY due to increase in sales volume
- Q1'FY26 EBITDA increased YoY, EBITDA margins lower QoQ due to annual maintenance shutdown at Spokane facility

CDMO - Sterile Injectables Vision 2030 : Double revenues by doubling of capacity at Spokane



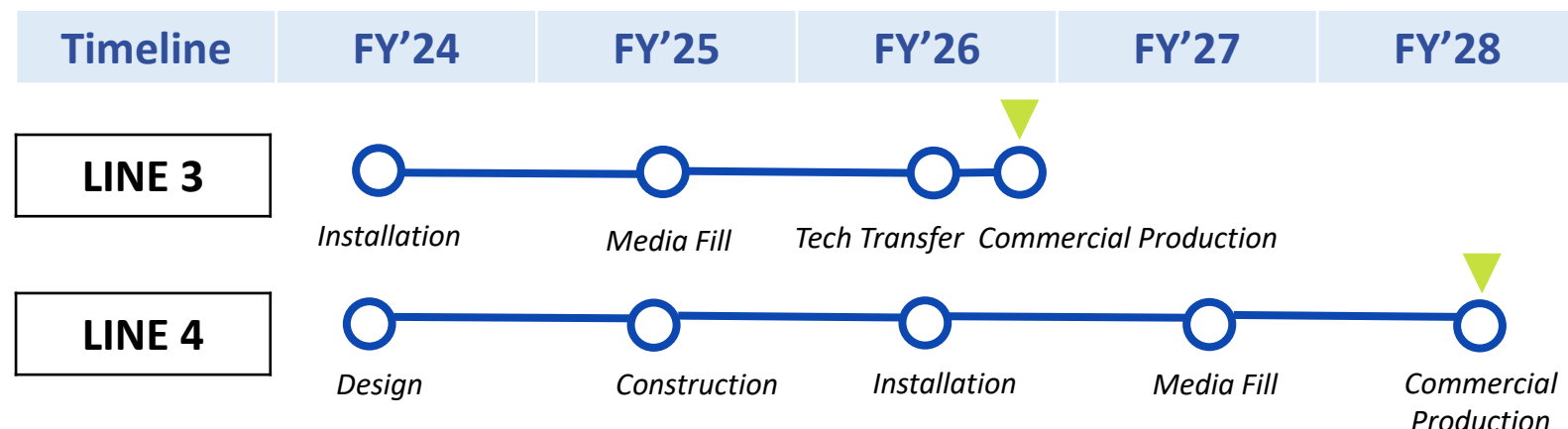
Line 3 to start commercial production in FY26

Multiple Tech transfers underway



Growth driver:

- Doubling Capacity



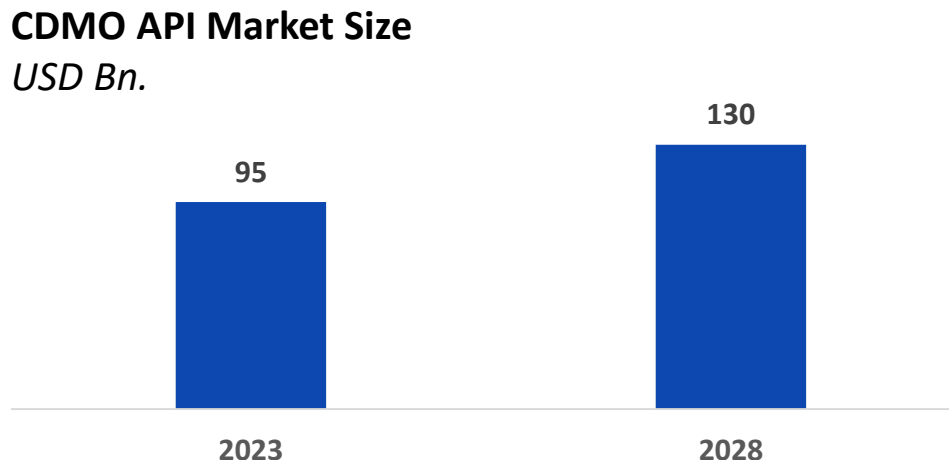
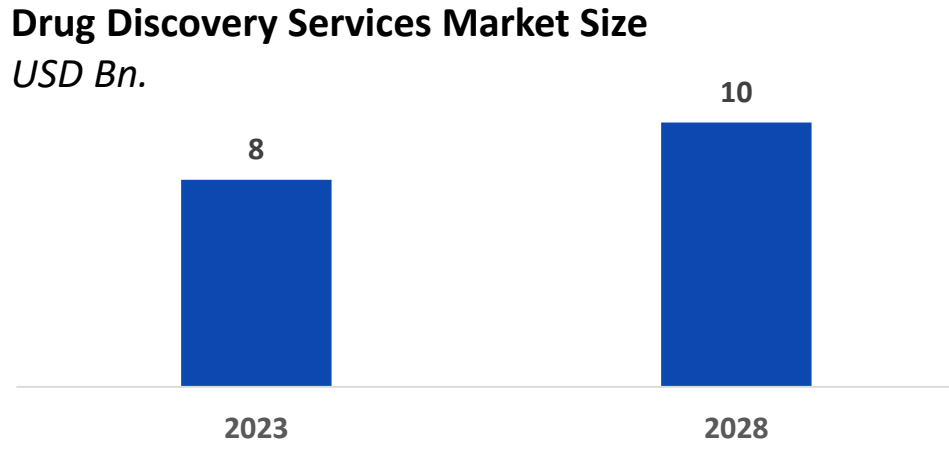
- Total investment at USD 285 Mn. (RoCE > 20%) including US Govt. funding of USD 150 Mn.
- New lines have incremental revenue potential of USD 160 Mn. to USD 180 Mn
- Excellent traction on RFPs incl. from Big Pharma. Expect to finalize these within FY26
- Expect to reach full Capacity utilization for Line 3 in 3 years vs 4 years (as expected earlier)

CRDMO: Drug Discovery Services, CDMO API



CRDMO: Drug Discovery, CDMO - API

India uniquely positioned to benefit from Friendshoring



Growth Drivers & Trends

Drug Discovery Market

- Biosecure Act advantage
- Rise in specialized technologies such as ADCs and oligonucleotides

CDMO API Market

- Rising interest in custom generics
- Rapid momentum in specialized CDMO services

We are a leading CRDMO for science with superior customer relationships



- **8 of the top 20 pharma** companies as customers with 5x increase in revenue share from Large Pharma
- **Indian Leader for “Integrated Drug Discovery”**, with a track record of +85 programs and Big pharma strategic partnership
- **Strengthen European penetration**, with multifold revenue increase
- **Fully integrated Chemistry powerhouse** from mg to multi-tons
- **Successful launch of new CDMO services** for Biotech and Large Pharma

...with state of the art integrated CRDMO platform



Drug Discovery Services & Early CDMO



**CoE Biologics
(St. Julien, France)**



**Integrated
Drug Discovery Centre
(IDDC, Bengaluru)**



**Chemistry Research
Innovation Centre
(CIRC, G. Noida)**



**Contract Development &
Manufacturing Centre
(API CDMC)**



**Advanced Intermediate
&
API Manufacturing**

~ 35 Scientists

Antibody Drug
Conjugates, Biologics

~ 350 Scientists

Identifying target to
candidate selection

~ 750 Scientists

Synthetic, Medicinal,
Analytical and
Computational Chemistry

~250 Scientists

Process Research Chemistry
& Manufacturing

900+ MT of capacity

US FDA, Japan PMDA,
Korea KFDA, Brazil ANVISA

**Immune - oncology
Expertise**

**+85
Integrated Programs
delivered**

**~40 clients
in last 3 years**

**From mg to kg
Supporting Scale-up up to
20 kg**

**Potent API expertise
OEB Class 1-4 API potency**

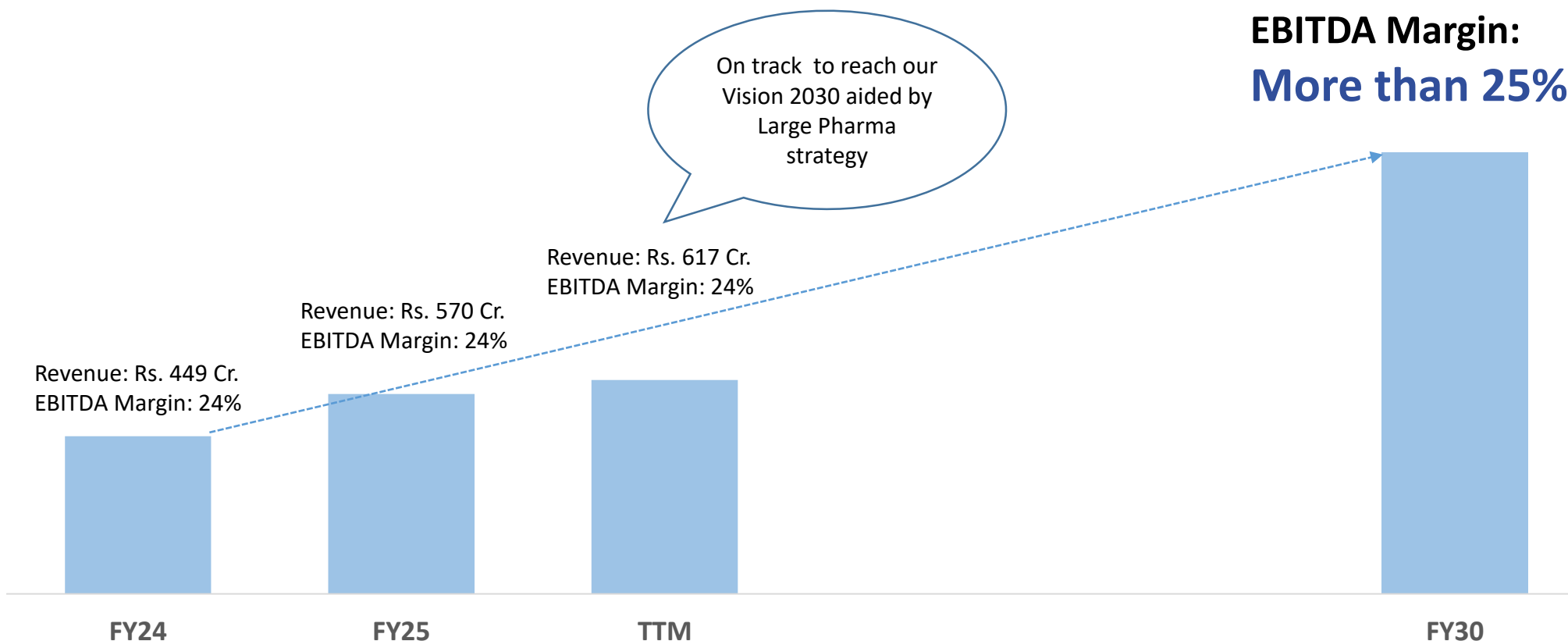
Drug Discovery Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	113	156	161	42%
EBITDA	22	41	32	46%
<i>EBITDA Margin (%)</i>	<i>19%</i>	<i>26%</i>	<i>20%</i>	<i>60 bps</i>

- Q1'FY26 revenue increased YoY from scaling Large Pharma contracts
- Q1'FY26 EBITDA increased YoY, EBITDA margins lower QoQ due to change in project mix and investment in business development

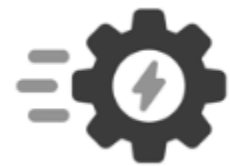
Drug Discovery Vision 2030 : Triple revenues & maintain profitability



Revenue:
~ 3x of FY24
EBITDA Margin:
More than 25%

Growth driver:

- Add Large Pharma



Proposed Biosecure Act

- Act passed in Sep'24 by US House of Representatives
- American pharma companies to look for alternatives besides China

- Executing strategy on Large Pharma
- Footprint in EU
- Introduction of ADCs, mAbs, and Biologics platforms

Drug Discovery Services: Expansion at current and new sites to enable revenue growth



Expansion at current sites, Greater Noida & Bengaluru



Expansion at new site, Devanahalli, Bengaluru



Capacity : 1,000 FTE's (FY25) → 2,000 FTE's (FY28) → 4,000 FTE's (FY30)

Increasing capacity in a phased manner ; Total Capex USD 150 Mn. (Expect RoCE > 20%)

Drug Discovery Services: Added capability in Biologics through strategic partnership with Pierre Fabre



- Expanded TAM by USD 1.4 Bn. in mAbs and ADCs
- Added strategic footprint in the EU
- Enhanced domain expertise in ADC
- Unique & cost-effective delivery model

Integration in progress; Investing in Business development team

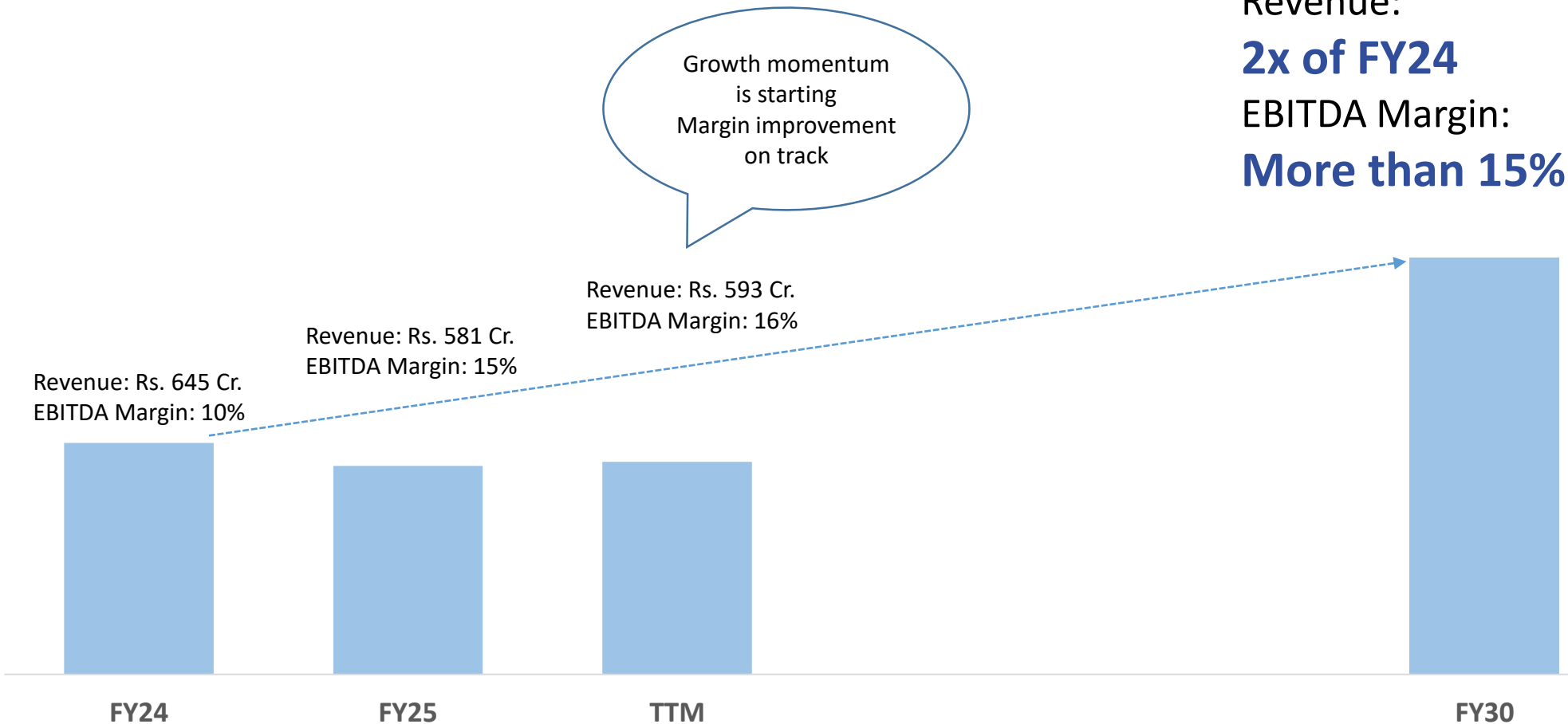
API Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	130	182	141	9%
EBITDA	16	39	22	36%
EBITDA Margin (%)	12%	21%	15%	310 bps

- Q1'Y26 revenue increased YoY on the back of increased volume in select products. Industry wide pricing pressure continues
- Q1'FY26 EBITDA margins increased YoY due to profitable product mix added by sustainable cost control measures

API Vision 2030 : Double revenues and increase profitability



Grow CDMO and custom manufacturing in API

Growth driver:

- Grow CDMO API



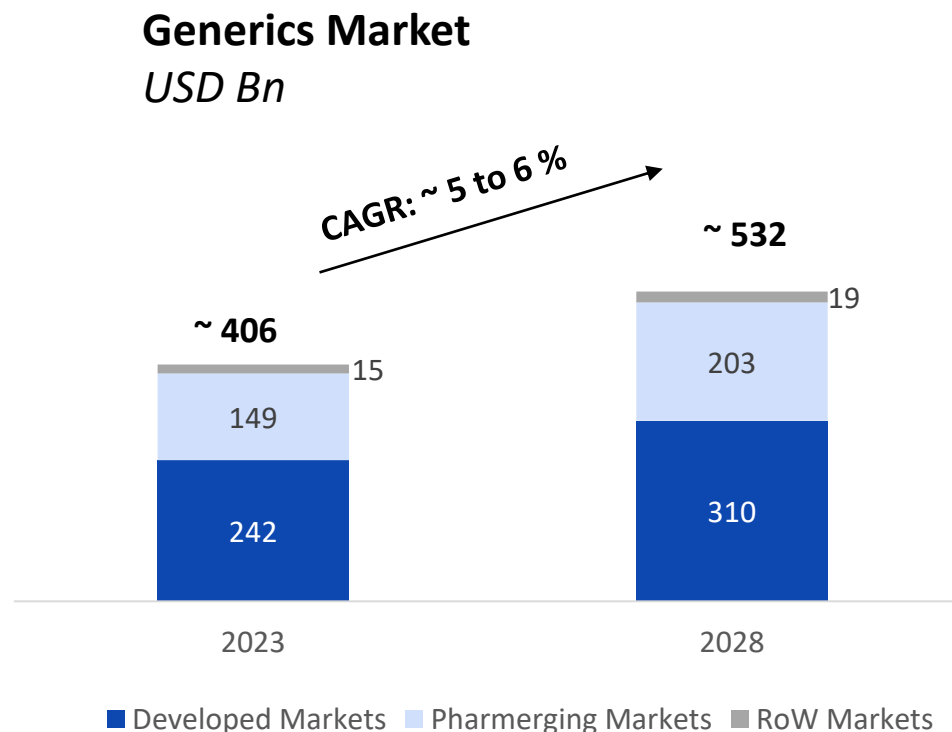
- **Further Strengthen CDMO:** Leverage GMP manufacturing capabilities for Innovative New Chemical Entities
- **Custom Manufacturing:** Partner with large pharma to manufacture products requiring life cycle management
- **China plus one strategy:** Resilient supply chain through increased backward integration & diversified supplier base

- **Proposed sale and transfer of API business to “Jubilant Biosys”, wholly owned subsidiary of company**
- Transaction will lead to **housing of Drug Discovery services** and **CDMO API business in one entity**
- Combined platform to **improve operational efficiency** and **superior brand recall of “Jubilant Biosys”**
- **Increase asset utilization of API business by improving revenue mix towards Custom manufacturing & CDMO**



Generics

Global Generics market expected to grow by ~ 5% to 6%



Growth Drivers and Trends

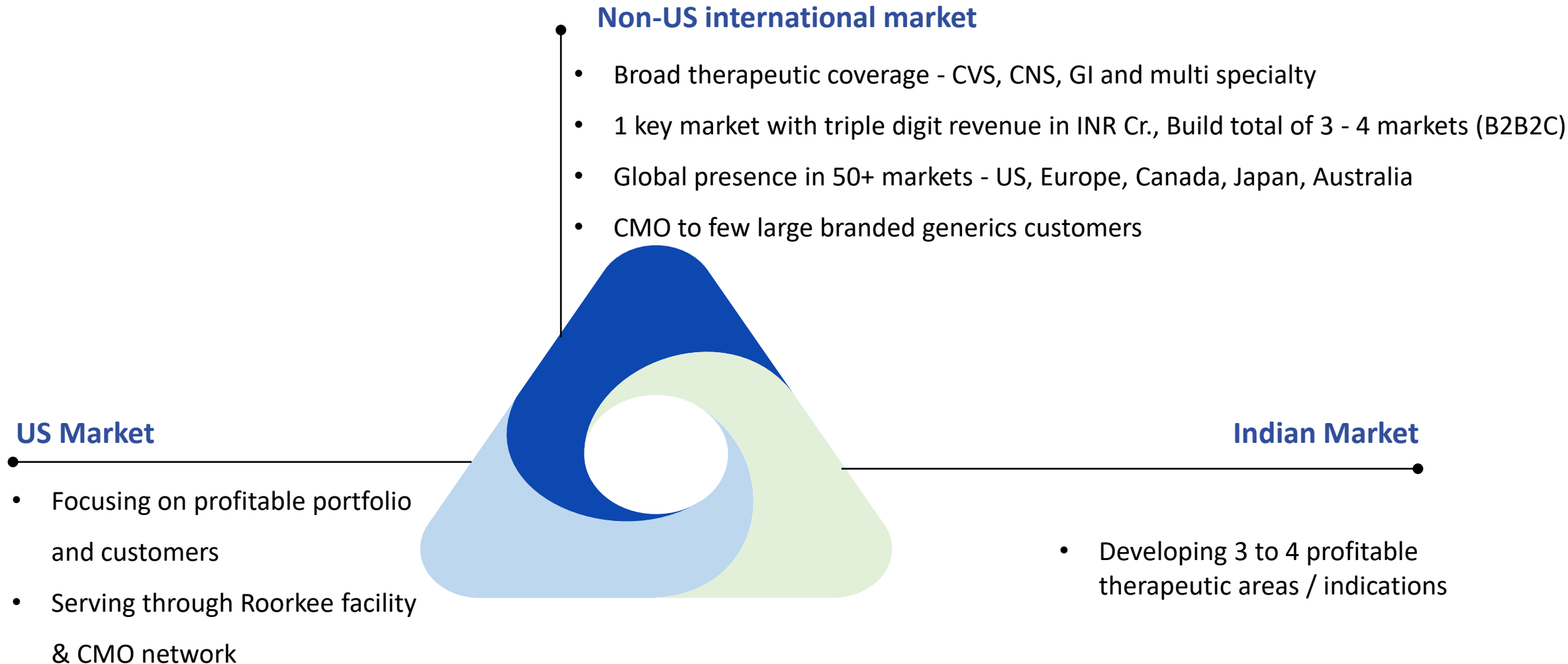
Developed Market

- US market to grow at 2%, signs of decrease in price reductions
- Non-US market to grow by 5 - 7%

India Market

- India market to grow in excess of 8%
- Brand building, in-clinic effectiveness of sales is key

We are building a growing, profitable & agile business model



Generics Financials : Q1'FY26



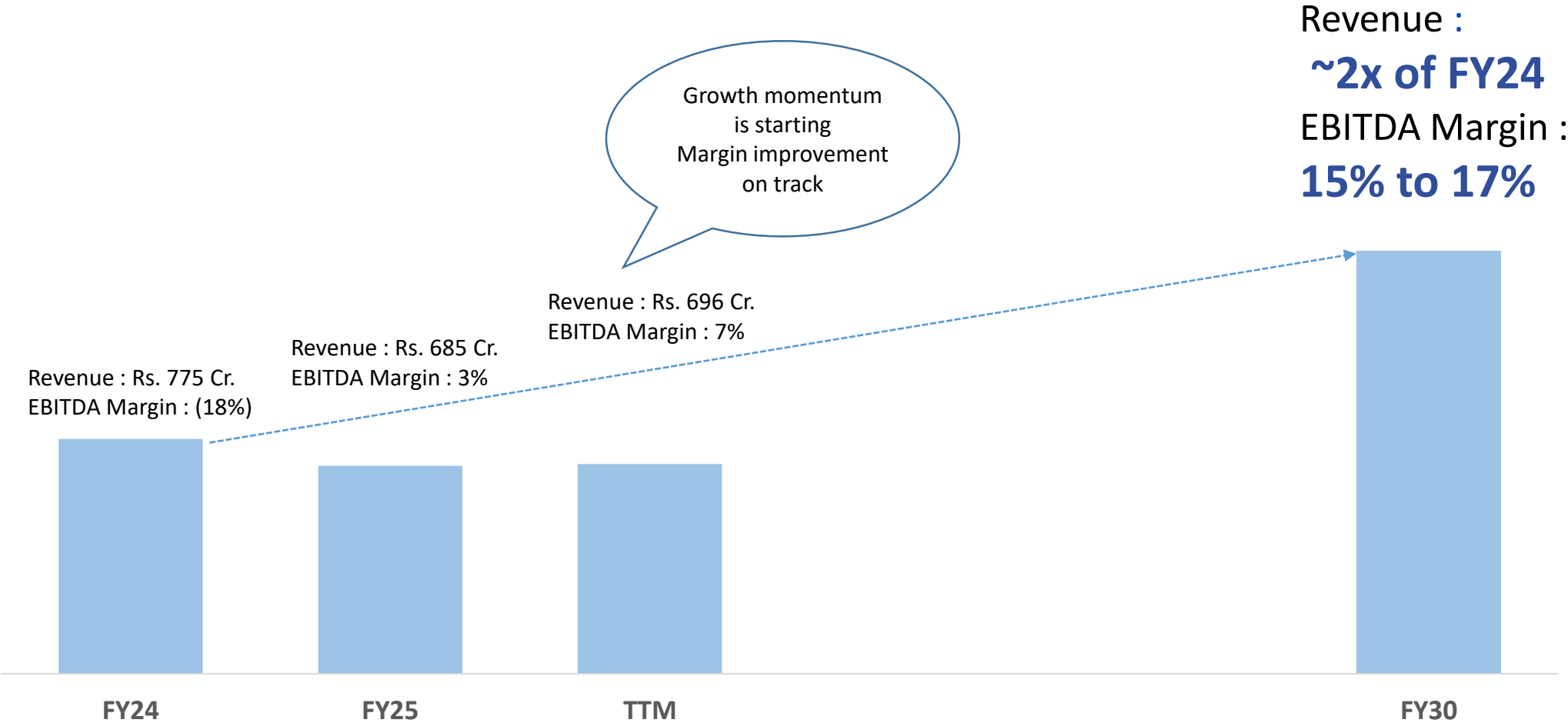
Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	156	157	166	7%
EBITDA	(11)	(17)	12	214%
EBITDA Margin (%)	(7%)	(11%)	7%	1,400 bps

- Q1'FY26 revenue increased YoY due to increase in revenue from Non-US markets
- Q1'FY26 EBITDA and EBITDA margins increased YoY due to focus on profitable products

Generics Vision 2030: Reach top quartile profitability for similar size companies



JUBILANT
PHARMOVA



Generics Growth Drivers



Launch new products

- Relaunch dormant ANDAs from Roorkee and CMO network
- Secure ANDAs approvals



Grow the profitable Non-US international market

- Launch 6 to 8 new products every year
- Scale 3 to 4 key markets



Build branded business

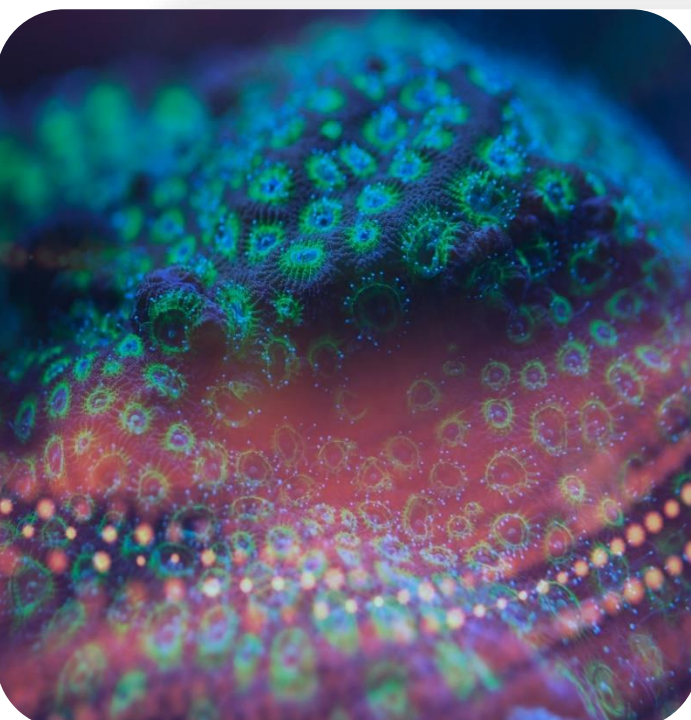
- Build presence in Diabetes, Dyslipidemia and Hypertension
- Scale in weight management
- Grow 1.5 times the Industry growth rate



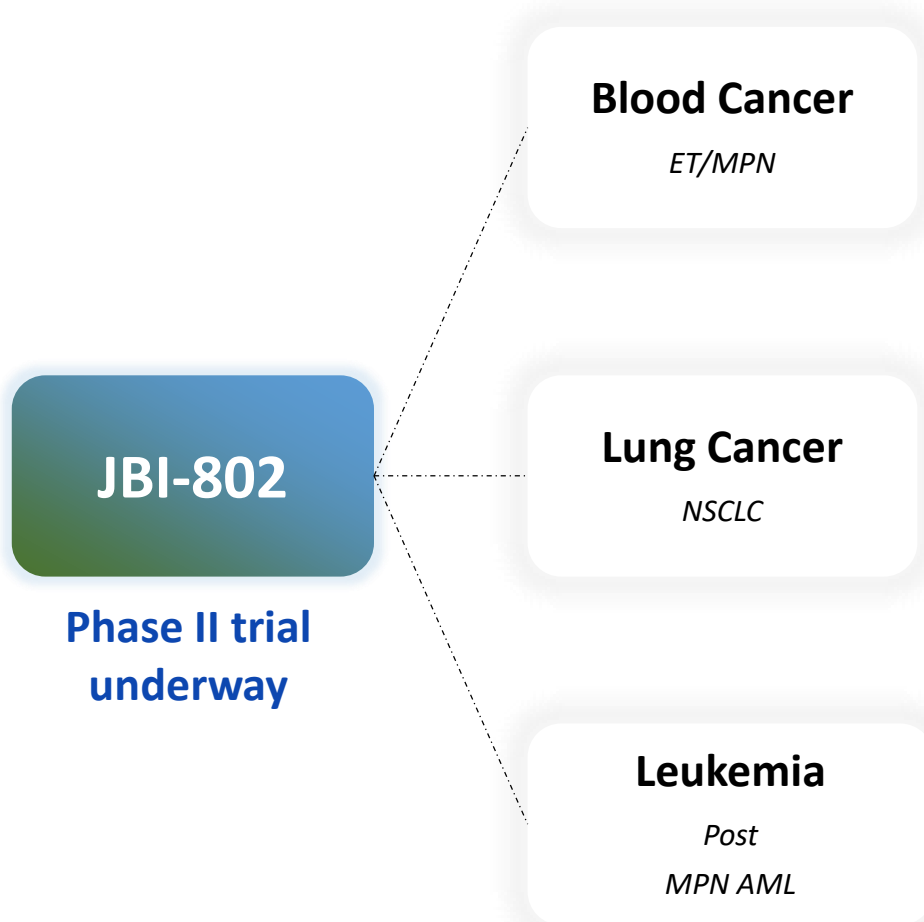
Proprietary Novel Drugs

Proprietary Novel Drugs



- 
- **Develop precision oral medicines** with enhanced safety and therapeutic efficacy
 - **Focused on specific set of patients**, not responding to other therapies
 - **Low-cost in-house discovery engine** to generate drug candidates, validated through partnerships
 - **Guided by world's leading oncologists** from Memorial Sloan Kettering and Dana Farber
 - **FDA Orphan drug designations** for leading programs JBI-802 and JBI-778

JBI-802 to address unmet medical needs in difficult to treat cancers



- **Company sponsored Phase II trial underway**
 - Highly differentiated for safety and efficacy than peers
 - Total Addressable Market in US: USD 3.3 Bn.
-
- **Investigator led trial initiated**
 - Demonstrated clinical efficacy in two NSCLC patients in phase 1 study
 - Total Addressable Market in US: USD 3.1 Bn.
-
- **Investigator led trial under planning**
 - Blood cancer progression to Leukemia is a serious complication
 - Total Addressable Market in US: USD 0.8 Bn.

JBI -802 has demonstrated transformative treatment in two patients

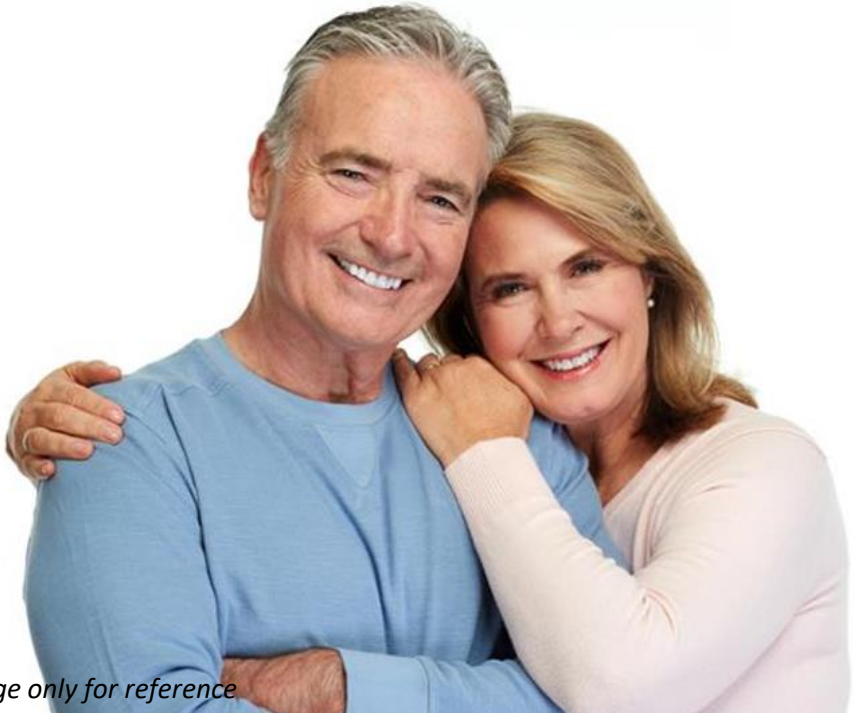
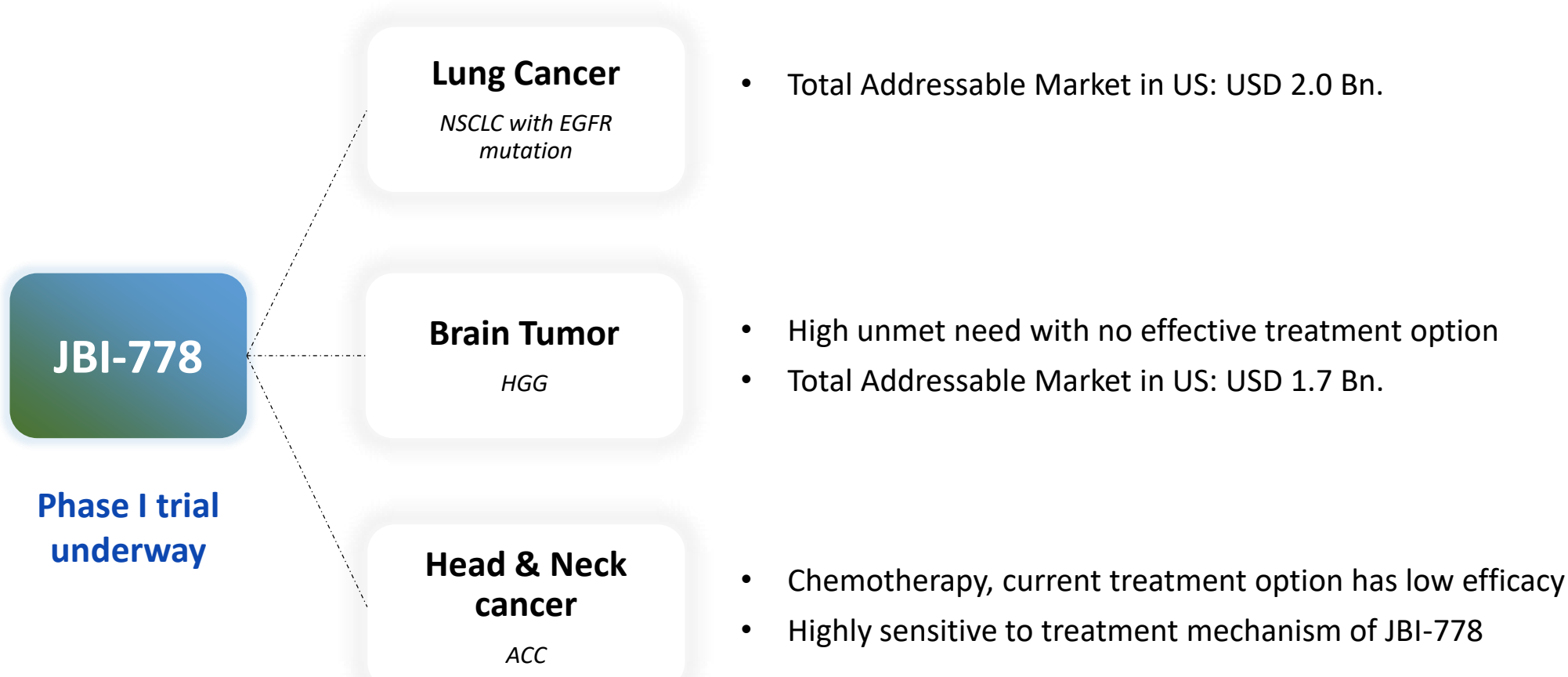


Image only for reference

- Non small cell lung cancer patient progressed to last stage after immunotherapy. Post taking JBI-802 treatment, patient has been doing very well even after two years. Major symptoms have disappeared with confirmed partial response with **~40% tumor reduction**
- **Over 50% shrinkage of the patient's liver metastasis** and a complete resolution of related portal hypertension and improvement in quality of life

JBI-778 to address unmet medical needs in difficult to treat cancers



Company sponsored First-in- human Phase I trial ongoing in India

Proprietary Novel Drugs Financials : Q1'FY26



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	0	0	0	
EBITDA	(6)	(4)	(6)	(3%)

- Continue to invest in a calibrated manner in two lead programs

Proprietary Novel Drugs to explore monetization



- Expect clinical data readouts in CY 2026
- **Explore monetization through licensing or external fund raising**

Consolidated Reported Financials – Q1'FY26

Solid revenue growth (YoY) along with EBITDA & PAT margin expansion (YoY)



Particulars (Rs. Cr.)	Q1'FY25	Q4'FY25	Q1'FY26	Y-o-Y
Revenue	1,732	1,929	1,901	10%
Other Income	14	12	12	
Total Income	1,746	1,941	1,913	10%
EBITDA	266	357	302	14%
<i>EBITDA Margin (%)</i>	<i>15.2%</i>	<i>18.4%</i>	<i>15.8%</i>	<i>60 bps</i>
Exceptional Income / (expense)	396	(3)	0	
PBT	500	206	154	
PBT Margin	28.6%	10.6%	8.1%	
Normalised PBT¹	104	209	154	49%
Normalised PBT Margin	5.9%	10.8%	8.1%	210 bps
Reported PAT	482	151	103	
Reported PAT Margin	27.6%	7.8%	5.4%	
Normalised PAT¹	69	139	103	48%
Normalised PAT Margin	4.0%	7.1%	5.4%	140 bps

Normalised PBT is after adjusting for Exceptional items

Normalised PAT is after adjusting for Exceptional items and tax

- Q1'FY26 **Revenue grew YoY** on the back of growth in revenue across all business units
- Q1'FY26 **EBITDA margins increased YoY** due to improved performance in CRDMO and Generics
- Q1'FY26 **Normalised PAT margins increased YoY** due to improved operating performance and reduction in finance cost

Key Ratios

Net Debt / Ebitda to remain range bound



Particulars (Rs. Cr.)	Mar 31, 2025	June 30, 2025
Net Debt (On constant currency, Net of DIC)	1,348	1,535
Net Debt / Equity	0.22	0.24
Net Debt / EBITDA (TTM)	1.1	1.2
Long Term Capex Creditors	453	455

- Investing consciously to maintain Net debt / Ebitda range bound

Sustainability

Received KPMG ESG Excellence award for Mid / Small Cap category in the Pharma & Healthcare in FY25



S&P Global



DJSI Score 60%



EcoVadis Score 65

KPMG

KPMG ESG Conclave and Awards 2024

Winner – Mid/Small Cap Category



B - Water Security, D - Climate



COMMUNITY MEMBER

2025

Member since 2005



CoP submitted,
3rd July 25

Member since 2010

All Indian manufacturing plants are
Zero Liquid Discharge

13 percentage points increase in renewable in
purchased power v/s FY 25 target

Sustainability Performance

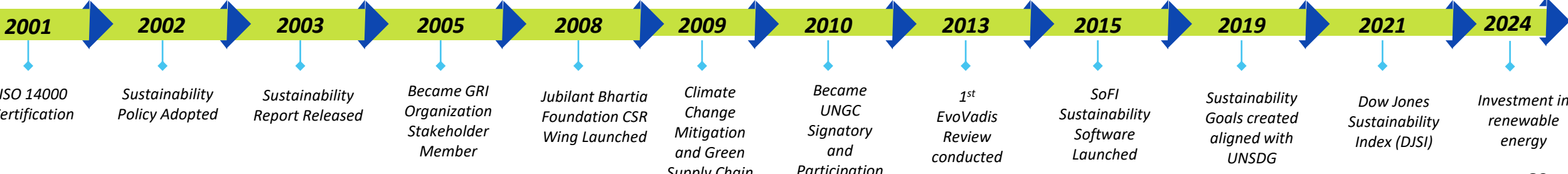
21% reduction in Specific Greenhouse
Gas Emissions v/s FY 25 target

19% reduction in Specific Water
Consumption v/s FY 25 target



Category I ERP

ESG Score 68



Summary – Q1'FY26



1

Radio Pharmaceuticals : Ruby-Fill® maintaining **growth momentum**. New Products to drive margin expansion
Radio Pharmacies : Competitive intensity higher in SPECT, **Commercial distribution of PLYARIFY® in PET** continue to grow

2

Allergy Immunotherapy : Revenue grew YoY; **EBITDA margins in normalized range**

3

CDMO Sterile Injectable : **Capacity expansion** at Spokane **on track**. Line 3 progressing ahead of expectations

4

CRDMO DDS: Continue to increase revenue share from large pharma clients. **Medium term outlook continues to be positive**
CRDMO API : Focus on profitable products and CDMO. **Taking initiatives to reduce operating costs**

5

Generics : Improving **growth & profitability outlook**

6

Prop Novel Drugs : **Patient dosing** progressing in both lead programs

Financial Results Table

Total Income (Rs. Cr.)	Q1'FY25		Q4'FY25		Q1'FY26	
Revenue (A)	1,732		1,929		1,901	
a. Radiopharma	832		895		869	
<i>Radiopharmaceuticals</i>	262		296		271	
<i>Radiopharmacies</i>	570		600		598	
b. Allergy Immunotherapy	168		192		181	
c. CDMO Sterile Injectables	324		340		370	
d. CRDMO	243		338		302	
<i>Drug Discovery Services</i>	113		156		161	
<i>CDMO – API</i>	130		182		141	
e. Generics	156		157		166	
f. Proprietary Novel Drugs	0		0		0	
<i>Unallocable Corporate Income</i>	10		7		13	
Other Income (B)	14		12		12	
Total Income (A+B)	1,746		1,941		1,913	
EBITDA (Rs. Cr.)	Q1'FY25	Margin	Q4'FY25	Margin	Q1'FY26	Margin
a. Radiopharma	138	17%	141	16%	136	16%
<i>Radiopharmaceuticals</i>	126	48%	136	46%	126	46%
<i>Radiopharmacies</i>	13	2%	6	1%	10	2%
b. Allergy Immunotherapy	63	38%	88	46%	63	35%
c. CDMO Sterile Injectables	57	18%	95	28%	62	17%
d. CRDMO	38	16%	79	23%	54	18%
<i>Drug Discovery Services</i>	22	19%	41	26%	32	20%
<i>CDMO – API</i>	16	12%	39	21%	22	15%
e. Generics	(11)	(7%)	(17)	(11%)	12	7%
f. Proprietary Novel Drugs	(6)		(4)		(6)	
<i>Unallocable Corporate (Expenses) / Income</i>	(15)		(26)		(18)	
Total EBITDA	266	15.2%	357	18.4%	302	15.8%

Revenue

Reach **2X** *from FY24 to FY30*

EBITDA Margin

23% to 25% *by FY30*

Net Debt

Zero *by FY30*

RoCE

High Teens *by FY30*

Annexure

Executive Leadership Team



Shyam S Bhartia
Chairman



Hari S Bhartia
Co-Chairman



Priyavrat Bhartia
Managing Director



Arjun S Bhartia
Joint Managing Director



Arvind Chokhany
Group CFO, Whole-time Director



Shantanu Jha
Group CHRO



Dr Tushar Gupta
Head - Corporate Strategy

Executive Leadership Team



Harsher Singh
CEO - Jubilant Radiopharma



Chris Preti
CEO - CDMO Sterile Injectables



Giuliano Perfetti
CEO - CRDMO, Biosys



Dr. Jaidev Rajpal
CEO - Jubilant Generics



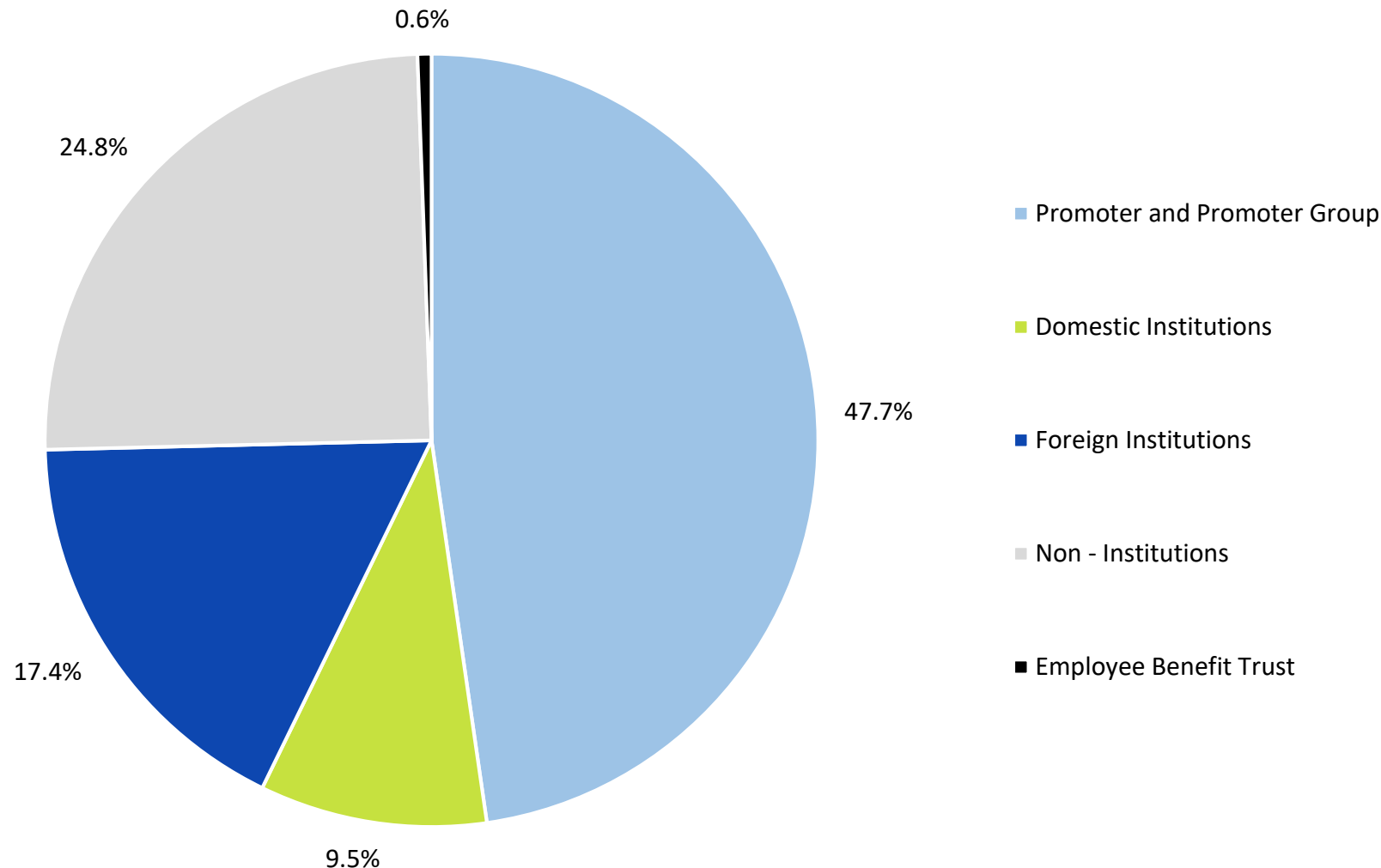
Kyle Ferguson
CEO - Allergy Immunotherapy



Dr. Syed Kazmi
CEO - Jubilant Therapeutics

Shareholding Pattern

As on 30th June 2025



Glossary



Abbreviation	Details
CVS	Cardiovascular System
CNS	Central Nervous System
CDMO	Contract Development Manufacturing Organization
CRDMO	Contract Research & Development Manufacturing Organization
F18	Fluorine-18 Radioisotope
PSMA	Prostate Specific Membrane Antigen
Lu177	Lutetium-177 Radioisotope
Ac225	Actinium-225 Radioisotope
MAA	Macro Aggregated Albumin
DTPA	Diethylenetriaminepentacetic Acid-Chelating Agent
HICON	Pharmaceutical Grade Radioactive Iodine
I 131	Iodine-131 Radioisotope
MIBG	Metaiodobenzylguanidine
USP (USP 825 Guideline)	U.S. Pharmacopeia (USP) general chapter ,825 (Related to Radiopharmaceuticals: Preparation, Compounding, Dispensing, and Repackaging)
Ga 68	Gallium-68 Radioisotope
Rb	Rubidium (chemical element)
Sr	Strontium (chemical element)
Cu 64	Copper-64 Radioisotope
NRC	Nuclear Regulatory Commission (U.S.)
GPOs	Group Purchasing Organisation
IDNs	Integrated Delivery Network
SCIL	Sublingual immunotherapy (Allergy treatment - Dust mites & Seasonal allergy)
SCIT	Subcutaneous Immunotherapy (Allergy treatment Insect venom, pet dander, Mold, and other allergens)
APAC	Asia Pacific
MEA	Middle East Africa
NSCLC	Non-small cell lung cancer
SCLC	Small cell lung cancer

Abbreviation	Details
MEA	Middle East Africa
LATAM	Latin America
LOE	Loss of exclusivity
FDA (US)	U.S. Food and Drug Administration
PMDA (Japan)	Pharmaceutical and Medical Device Agency
KFDA (Korea)	Korea Food Development Authority
ANVISA (Brazil)	Brazilian Health Regulatory Agency
TGA (Australia)	Therapeutic Goods Administration
API	Active Pharmaceutical Ingredient
MENA	Middle East North Africa
GMP	Good Manufacturing Practices
B2B2C	Business-to-Business-to-Consumer
B2B	Business-to-Business
ET/MPN	Essential thrombocythemia / Myeloproliferative neoplasm (rare chronic blood cancer)
coreST	CRISPR-Cas9 Endomorphic RNA Symptomatic Inhibitor (RNA based therapy targeting genetic disease)
Inhibitor/ Epigenetic Modulating Agent	Medications that modify gene expression patterns
PRMT5 Inhibitor	Protein Arginine Methyltransferase 5 inhibitor (Blocks enzyme activity involved in adding methyl groups to arginine residues, affecting gene expression regulation)
Brain Penetrant	Cerebral blood flow enhancers or cognitive-enhancing drugs (supplements)
PD-L1 Inhibitor	Programmed death Ligand-1 inhibitor (blocks the PD-L1 pathway, enhancing immune response against cancer cells)
PAD4 Inhibitor	poly(ADP-ribose) polymerase 4 inhibitor (Disrupts DNA repair mechanisms in cancer cells, leading to their death)
LSD1/HDAC6 inhibitor	Lysine specific demethylase 1/Histone deacetylase 6 inhibitor (Blocks enzymes involved in modifying histones, impacting gene expression regulation in cancer therapy)
NSCLC	Non-small cell lung cancer
SCLC	Small cell lung cancer

A tray filled with numerous test tubes containing various colored liquids, including blue, green, yellow, and orange.

Thanks!



Q1'FY26 Q&A

Disclaimer

Statements in this document relating to future status, events, or circumstances, including but not limited to statements about plans and objectives, the progress and results of research and development, potential product characteristics and uses, product sales potential and target dates for product launch are forward-looking statements based on estimates and the anticipated effects of future events on current and developing circumstances. Such statements are subject to numerous risks and uncertainties and are not necessarily predictive of future results. Actual results may differ materially from those anticipated in the forward-looking statements. Jubilant Pharmova may, from time to time, make additional written and oral forward looking statements, including statements contained in the company's filings with the regulatory bodies and its reports to shareholders. The company assumes no obligation to update forward-looking statements to reflect actual results, changed assumptions or other factors.

Radiopharmaceuticals

Q1. Can you talk about growth in Ruby-Fill®?

Answer: Ruby-fill® is a best in class Positron Emission Tomography (PET) radiopharmaceutical product used for Cardiac imaging through a non-invasive procedure of the myocardium, to evaluate regional myocardial perfusion in adult patients with suspected or existing coronary artery disease. Our product is superior due to longer shelf life leading to more scans per generator, better and consistent image quality due to the patented saline push feature and multiple safety features.

We have witnessed strong installations in Q1'FY26. Our focus now is value engineering to improve margin and increase consistency and we will be deploying in a short period an AI enabled 3D cardiac blood flow quantification system that allows to get an image and deliver it under 75 seconds through artificial intelligence.

Q2. Can you talk about the sales of SPECT product portfolio in Q1'FY26?

Answer: We continue to maintain strong position in our SPECT portfolio. We are happy with the offtake of our new products.

We have seen a generic entry in DTPA in the US market. We expect loss of market share in DTPA from the current year. To counter the same, we are working to launch new products. We expect to file one new product in FY26.

Q3. What is the timeline of the MIBG Launch? What is the patient recruitment status & expected date of result?

Answer: MIBG is targeting paediatric patients with high-risk Neuroblastoma. Neuroblastoma is a type of cancer that forms in certain types of nerve tissues. The incidence of Neuroblastoma in the US market is estimated at 800 new cases per year and the relapse / refractory cases are estimated at 400 per year.

MIBG clinical trials are progressing well. We have completed the dosing of Phase two trials. We plan to send the data package to FDA by H2'FY26 and the followed by a pre NDA meeting, we shall file for approval. We expect to launch MIBG by FY27 after securing product and manufacturing approval.

Q4. Can you give us some more colour on the product pipeline?

Answer: We have a very strong pipeline of products in PET and SPECT with an Addressable Market at approx. USD 550 million. These products are generics or 505 B(2) versions of existing products and will be launched from FY27 to FY29. In addition to that, on the therapeutics side, we are working on MIBG.

Q5. Can you explain Q1'FY26 Radiopharmaceutical results?

Answer: Q1'FY26 revenue grew 3% YoY to Rs. 271 Cr. on the back of growth in Ruby-Fill®. Q1'FY26 EBITDA remained stable YoY at Rs. 126 Cr.

Radiopharmacy

Q6. Can you talk about Industry demand? Where are we in the execution of new PET Radiopharmacy project?

Answer: The PET Imaging market is growing rapidly on the back of new products. There are multiple commercial products today for Prostate Cancer, Alzheimer's, Breast Cancer, Parkinson Disease, Cardiac Imaging and others. There are many products in the pipeline, which shall be commercialised in the coming years.

We are pleased to share that we have forged multiple partnerships with Radiopharmaceutical manufacturers in the PET imaging. Notable ones include Life Molecular Imaging's F18 Neuraceq and Lantheus' F18 Pylarify. We expect PET revenue mix to increase in the back of increase sales of PYLARIFY®, which is an industry leading prostate cancer diagnostic imaging agent.

We also announced USD 50 million investment to expand our PET radiopharmacy network from three (3) to nine (9) sites and therefore positioning us to secure long-term contracts with the leading PET radiopharmaceutical manufacturers. These new PET radiopharmacies shall be fully operational in FY28 and shall start to contribute significantly to the top line and bottom line. We expect a RoCE in excess of 20% on our investment.

Q7. Can you explain Q1'FY26 Radiopharmacy results?

Answer: Q1'FY26 revenue grew 5% YoY to Rs. 598 Cr. on the back of increase in volume from new PET products. Q1'FY26 EBITDA stands at Rs. 10 Cr., lower YoY due to increase in competitive intensity in SPECT radiopharmacies. Revenue from our current PET radiopharmacies continue to increase.

Allergy Immunotherapy

Q8. What are the growth levers in this business?

Answer: The business is moving ahead on a three-pronged growth strategy.

The first is to strengthen the existing position in both Venom and Non-Venom segment in the US. We are increasing customer awareness about the importance of bee sting allergy treatments through targeted marketing campaigns. We are also working to increase revenue in the US allergenic extract market through an emphasis on science and product differentiation.

The second is to expand its footprint in select international markets, through strategic partnerships and an expanded distribution channel.

Last and most important is to develop new products and technologies by increasing investment in R&D. The business continues to develop innovative products to address various allergies, as evidenced by the 2023 launch of Ultra filtered Dog Hair and Dander extract. This product provides optimal treatment, ensuring dependable consistent results, and efficacious dosing without precipitate formation.

Q9. Can you explain Q1'FY26 Allergy immunotherapy results?

Answer: In Q1'FY26, Revenues grew by 8% on YoY basis to Rs. 181 Cr. on the back of growth in revenue from the US market. EBITDA remained stable YoY at Rs. 63 Cr. EBITDA margin for Q1'FY26 stands at 35%. EBITDA margins are lower QoQ as Q4'FY25 margins were exceptionally higher due to high production volume.

CDMO Sterile Injectable

Q10. Can you talk about the overall demand scenario in the sterile fill and finish market?

Answer: The global CDMO Sterile fill and finish market is expected to grow from USD 13 billion in 2023 to USD 20 billion by 2027 at a CAGR of 11%. The demand drivers include an increasing number of injectables in the development pipeline driven by biologics, predominantly in Vial format and an increase in loss of exclusivities.

As big pharma is focused on internalising only select capabilities, this increase in demand is being outsourced to specialised CDMO Sterile fill and finish companies. In addition, there is a strong Customer preference for on-shore capacity due to higher-value products, regulatory & supply chain advantages.

The drug shortages in the injectables in the US are signalling that the demand still runs higher than the supply of the capacity and therefore there is a need of significant new capacity. In particular, a recent McKinsey report highlighted a 6.8Bn demand vs. 6.1Bn supply specifically for sterile vials – a 700 Mn. sterile vial shortfall.

The proposed Biosecure Act and the Catalent acquisition by Novo Holding shall further widen the gap between demand and supply in the US.

In addition to that, the large innovator pharma companies, for their US requirements, are now looking to create an alternate manufacturing site in the US as a risk management measure in the event of tariff imposed by the US Govt.

Q11. What is the project status of the expansion project in Spokane? What is the maximum revenue potential for Line 3 & 4 and how much time it shall take to fill up the lines?

Answer: The capacity expansion program in Spokane, Washington, USA is on track. We are pleased to share that we have successfully completed Media-Fills at Line 3 along with our first product qualification batches (PPQs). Multiple customer's technology transfer programs are underway. The commercial production at Line 3 is expected to start in the current financial year.

The revenue potential for Line 3 and Line 4 combined is estimated at USD 160 to 180 million. The large innovator pharma companies are now looking to create an alternate manufacturing site in the US as a risk management strategy to mitigate any potential tariff's imposed by the US govt. In light of that, we see an excellent traction in Requests for Proposals (RFPs) for Line 3 including from Big Pharma. We expect to finalise these within FY26. Therefore, we expect to reach full capacity utilisation for Line 3 in 3 years now vs 4 years, which was expected earlier. Also, we expect to clock higher than normalised EBITDA margins on the incremental revenue from Line 3 and Line 4 on the back of improved pricing due to newer technology and lower incremental overheads.

Q12. Can you give us an update on Montreal facility?

Answer: Our Montreal facility received OAI classification post FDA audit in FY25. Pursuant to the USFDA audit observations and in light of recently issued new and stringent cGMP guidelines for sterile fill and finish manufacturing facilities, we implemented corrective and Preventive actions (CAPA's) in our manufacturing set up at Montreal facility. Once the CAPA implementation was complete, we restarted sterile

injectables operations successfully in middle of Q3'FY25 and are stable now. We plan to return to production for ophthalmic line in H2'FY26.

Also at the Montreal facility, we have announced an investment of USD 114 million towards the expansion of our liquid and lyophilization sterile fill operations. Of the total investment, approximately USD 40 million of project cost will be funded through concessional loans from the Canadian Government and the balance from internal accruals.

Additionally, we are investing in the area of sterile ophthalmic by setting up a 200-bottle-per-minute plant at the Montreal, Canada facility given the high Requests for Proposals (RFPs) This ophthalmic line is currently undergoing validations. It is expected to be commercially qualified by the end of FY26.

Q13. Can you explain Q1'FY26 CDMO Sterile Injectables results?

Answer: Q1'FY26 revenue increased by 14% YoY at Rs. 370 Cr on the back of increase in sales volume. EBITDA increased by 9% to Rs. 62 Cr. on the back of revenue growth. EBITDA margins are lower QoQ on the back of annual maintenance shutdown at Spokane facility

CRDMO – Drug Discovery

Q14. We have seen an impressive revenue growth in Drug Discovery services? Can you talk about it the growth trajectory going forward?

Answer: We are bullish on the mid and long term prospects of the CRO industry in India due to talent availability & gradual shifting of demand due to the preference for “friend shoring” due to proposed Biosecure ACT. We are increasing our partnership with large Pharma companies doing, leveraging our infrastructure, capacity and capabilities expanded during last two years. As a testament, we on boarded three large pharmaceutical companies in last one year and we are scaling these contracts.

We are well prepared to further scale up Infrastructure and scientific talent to take advantage of the upcoming in CRO demand. We expect revenue growth to continue on the back of our large pharma strategy and the industry tailwinds.

We have talked about increasing our FTE capacity by four times to 4,000 FTE's in phased manner to cater to increasing demand.

Q15. Can you explain Q1'FY26 CRDMO Drug Discovery results?

Answer: In Q1'FY26, the Drug Discovery business revenue grew by 42% to Rs. 161 Cr and EBITDA grew by 46% to Rs. 32 Cr. Q1'FY26 revenue increased YoY due to increase in revenue from large Pharma customers. We have integrated new R&D facility in France and are now investing in business development. EBITDA Margins are lower on QoQ due to change in project mix and investment in business development.

CRDMO – API

Q16. What is the rationale of sale and transfer of API business to Jubilant Biosys?

Answer: The transaction will result in housing of the drug discovery business and CDMO API business in a single business entity. This combined platform will improve the operational efficiency in the business and lead to superior brand recall of "Jubilant Biosys" as provider of end-to-end CRDMO (Drug discovery, Early CDMO, late CDMO and commercial manufacturing) services by the large pharmaceutical & Biotech customers. The transaction will also help to improve asset utilisation of API business by improving the revenue mix towards Custom manufacturing & CDMO.

Q17. Can you explain Q1'FY26 CRDMO API results?

Answer: The API business revenue grew by 9% to Rs. 141 Cr in Q1'FY26. EBITDA grew by 36% to Rs. 22 Cr. EBITDA margins improved by 310 basis points due to profitable product mix. Industry wide pricing pressure still continues.

Generics

Q18. Can you explain Q1'FY26 generics results?

Answer: In Q1'FY26, the generics business revenue grew by 7% to Rs. 166 Cr. EBITDA for the period stands at Rs. 12 Cr. Revenue increase is primarily driven by Non US markets. Ebitda margins improved by 1400 basis points YoY due to focus on profitable products.

Of the last 4 quarters, Q2 and Q3'FY25 had positive EBITDA while the EBITDA for Q4 FY'25, was negative. The business returned to profitability in Q1'FY26. This, QoQ variability in margins is on the expected lines. On annual basis, however, the Generics business expects to achieve stability and move towards Generics Vision 2030 as shared earlier.

Q19. Can you tell us your plans for new product launches?

Answer: Since April'24, we secured approval of (9) ANDA's from our pipeline and acquired (2) ANDAs. We plan to launch six to eight products per annum in our US and non-US international markets in the current year. Therefore, we have an improving growth and profitability outlook.

In line with our communication, we are ramping up exports to the US markets from our Roorkee facility in a meaningful and gradual manner. We have also started supply of products from our Contract manufacturing partners to the US market in line with our plan.

Prop Novel Drugs

Q20. What is the status of clinical trials of your lead programs JBI-802 and JBI -778?

Answer: The Company's most advanced program (CoREST inhibitor) JBI-802 Phase 1 clinical data established safety and further, dose dependent platelet effect was seen in the clinic at higher doses, establishing application in Essential Thrombocythemia (ET) and other Myeloproliferative Neoplasms (MPNs). In light of these, we have started a phase II clinical trial to treat ET and MPN patients with thrombocytosis (high platelets).

The phase I trial also showed anti-tumour response in two lung cancer patients with a good safety profile. One non-small cell lung cancer (NSCLC) patient with STK11 mutations, having progressed on prior doublet immune-oncology (IO) therapy, showed a confirmed partial response (tumour reduction). Generally, the survival rate is very low in such cases, however, the patient has responded well to JBI-802 monotherapy with meaningful improvement in quality of life and continues to be on treatment with our novel oral pills for over years. Therefore, a larger investigator led clinical trial in NSCLC is being initiated at Christ Hospital in Ohio, USA.

The Company is also in discussions with Memorial Sloan Kettering for an investigator led trial in post MPN AML (Erythroleukemia).

The second program (PRMT5 inhibitor) is JBI-778, which is the next generation, small molecule, orally available and brain penetrant oral pill for select cancers. We have now, launched a phase I, first in human study, for this molecule, in patients with specific cancer sub-sets at major oncology centers in India.

Consolidated Financials

Q21. Can you talk about financial performance in Q1'FY26 ?

Answer: In Q1'FY26, Revenue grew by 10% on a YoY basis to Rs. 1,901 Cr. on the back of growth in revenue across all business units. EBITDA grew by 14% on a YoY basis to Rs. 302 Cr. due to improved performance in CRDMO and Generics. Q1'FY26 normalised PAT increased by 48% on a YoY basis to Rs. 103 Cr. on the back of improved operating performance and reduced finance cost.

Q22. What is the outlook for FY26?

Answer: In Q1'FY26, revenue grew by 10% on YoY basis. We expect growth momentum to further strengthen. In Q1'FY26, EBITDA margins expanded by 60 basis points on YoY basis. We expect EBITDA margins to strengthen on a full year basis as well. In Q1'FY26, Normalised PAT margins expanded by 140 basis points on YoY basis. We expect PAT margins to strengthen on a full year basis as well. We expect capex intensity to continue at same pace as in Q1'FY26 and hence Net debt/Ebitda to marginally increase in FY26 over FY25.

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