

August 10, 2026

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
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Sub: Press Release alongwith Earnings Presentation on the financials and operational performance of the Company for the quarter ended June 30, 2026

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, 2026.

The above - mentioned documents would 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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**Jubilant Pharmova Limited**

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PRESS RELEASE

Noida, Aug 10, 2026

JUBILANT PHARMOVA – Q1'FY27 RESULTS*On track towards Vision 2030**Solid Revenue growth across all business segments continues*

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	1,901	2,290	2,229	17%
Total Income	1,913	2,314	2,249	18%
EBITDA	302	363	268	(11%)
EBITDA Margin (%)	15.8%	15.7%	11.9%	(385) bps
Reported PAT	103	119	56	(45%)
Reported PAT Margin	5.4%	5.2%	2.5%	(285) bps

The Board of Jubilant Pharmova Limited met today to approve financial results for the quarter ended June 30, 2026.

Commenting on the Company's performance for Q1'FY27, **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. 2,229 Cr. for Q1'FY27, which reflects a solid growth of 17% on YoY basis. Revenue growth is broad based across all our business segments, but particularly strong in CDMO Sterile Injectables on the back of technology transfer revenues from the new & third line. EBITDA for the quarter stands at Rs. 268 Cr. EBITDA margins declined year-on-year, primarily due to the unavailability of high-margin SPECT radiopharmaceutical products in the Radiopharmaceuticals business. Additionally, margins were impacted by negligible third-party revenues and higher operating expenses, including incremental remediation costs at the CMO Montreal facility. Reported PAT for the period stands at Rs. 56 Cr. PAT margins decreased due to lower operating profitability and increase in depreciation.

During Q1'FY27, post successful media-fills, Commercial batch production for SPECT radiopharmaceuticals has started and these batches are expected to be released in Q2'FY27. By H2'FY27, all the SPECT Radiopharmaceutical products are expected to be available. Therefore, we anticipate EBITDA margins to start to expand from H2'FY27 onwards.

During Q1'FY27, we saw continued growth momentum in the Ruby-Fill® installs. In the Allergy Immunotherapy business, we witnessed increase in demand from both markets, US and Outside US. In the CDMO Sterile Injectables business, we saw incremental revenues from technology transfer programs, at Line 3 in Spokane. In the CRDMO business, we are witnessing increase in revenues from Biotech customer segment and custom manufacturing. In the Generics Business, we launched two new products. Lastly, in our Proprietary Novel drugs business, we continue to make progress in JBI-802 and JBI-778 clinical trials."



Segmental Business Performance

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

Radiopharmaceuticals Q1'FY27 revenue grew by 19% to Rs. 322 Cr. and EBITDA for the period stood at Rs. 110 Cr. EBITDA margins decreased YoY due to unavailability of certain SPECT products. At CMO Montreal, Post Successful media-fills, Commercial batch production for SPECT radiopharmaceuticals has started and the batches will be released in Q2'FY27. By H2'FY27, all the SPECT Radiopharmaceutical products are expected to be available. In the Ruby-Fill®, as we can demonstrate superior value proposition against competition, we continue to attract new channel partners. Our Ruby-Fill® install base has grown by 26% in Q1'FY27 on an annualised basis. This improved scale is also helping to increase EBITDA margins in this product category. We are on track to introduce multiple new products in the PET and SPECT imaging from FY28 to FY29. The dosing for Phase 2 clinical trial for MIBG is complete and we expect to do NDA filing by H2'FY27.

Radiopharmacy Q1'FY27 revenue grew by 17% YoY to Rs. 700 Cr. on the back of increase in volume from certain PET products. EBITDA for the period grew by 19% to Rs. 12 Cr. We continue to see high competitive intensity in the SPECT Radiopharmacy segment. On the other hand, we continue to see revenue growth and strong EBITDA margins in our PET manufacturing facilities (3 sites).

The proposed investment of US\$ 60+ million in six (6) new PET manufacturing network is underway. The new PET manufacturing sites will become operational in FY28. This investment will take the overall PET radiopharmacy network to Nine (9) sites, thereby strongly positioning Jubilant Pharmova's radiopharmacy network as the second largest in the US and shall drive the future business growth & profitability.

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

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

In Q1'FY27, revenues grew by 18% to Rs. 214 Cr., driven by strong growth in the US & outside US markets. EBITDA grew by 5% to Rs. 66 Cr. EBITDA margins reduced YoY due to lower production.

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

Q1'FY27 revenue grew by 34% to Rs. 496 Cr. due to incremental revenue from Line 3. EBITDA for the period stood at Rs. 45 Cr. EBITDA margins were lower YoY due to negligible third-party revenues & higher operating expenses including incremental remediation cost at Montreal facility.

At the Spokane facility, the capacity expansion program remains on track. Following the launch of our third Sterile Fill & Finish line (Line 3) in Q2'FY26, we are successfully ramping up revenues from technology transfer programs. Currently, 10+ products across multiple formats and vial sizes are undergoing technology transfer on Line 3. We also onboarded one of the world's largest oncology products on Line 3. Commercial batch production is expected to commence in late FY27, subject to FDA approval of these products. Revenue at Spokane grew by 43% to Rs. 494 Cr. and EBITDA grew by 49% to Rs. 117 Cr. EBITDA margins also expanded by 100 basis points to 24%.

Considering the new tariffs imposed by the US Government, large innovator pharmaceutical companies are increasingly seeking high-quality, US-based manufacturing, specifically, those with significant capacities with isolator technology. As a result, we are seeing strong traction in Requests for Proposals (RFPs) for the new lines.



We are happy to share the completion of installation of Line 4. Post media-fills, Line 4 is expected to start generating technology transfer revenues in Q4'FY27.

At Montreal facility, In Q1'FY27, we progressed to stabilize production for Radiopharmaceutical products. We also expect the operating losses to reduce from H2'FY27 due to increase in sales of SPECT Radiopharmaceutical products. Post the receipt of warning letter in Q1'FY27, we have responded to the FDA and apprised them on our remediation actions. In the medium term, we anticipate that the new isolator-based fill-and-finish line (Line 5) will start generating revenues from FY29 onwards, thereby supporting the site for a profitable growth.

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

In Q1'FY27, the Drug Discovery business revenue grew by 8% to Rs. 174 Cr. EBITDA for the period grew by 43% to Rs. 45 Cr. EBITDA margins expanded by 630 basis points to 26%. In the short term, we expect competitive intensity to increase in the large-pharma customer segment, while demand conditions in the biotech segment are expected to improve. In the medium term, we expect to deliver healthy revenue growth & steady margins.

In the API business, revenue for Q1'FY27 stood at Rs. 135 Cr. EBITDA for the period stood at Rs. 19 Cr. Revenue and EBITDA margins decreased YoY due to the industry wide pricing pressure. RM cost increase due to middle east conflict is expected to impact the margins in the subsequent quarters. We are consciously moving the revenue mix towards profitable products. Going forward, we expect the custom manufacturing revenue mix to drive the utilisation.

Generics – Building a growing, profitable & agile business model

In Q1'FY27, the Generics business revenue grew by 4% to Rs. 173 Cr on the back of launch of 2 new products. EBITDA for the period stood at Rs. 4 Cr. EBITDA margins decreased YoY due to change in product mix. Looking ahead, we are preparing to launch multiple products in FY27 to drive revenue growth & profitability.

Proprietary Novel Drugs – Innovative biopharmaceutical company developing breakthrough therapies

The global clinical trials for our lead programs, Phase I/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), Adenoid Cystic Carcinoma and high-grade Glioma are actively enrolling patients and progressing in line with our expectations.



About Jubilant Pharmova Limited

Jubilant Pharmova Limited is an integrated global pharmaceutical company engaged in Radiopharma, Allergy Immunotherapy, Contract Development and Manufacturing of sterile injectable, Contract Research Development and Manufacturing, Generics and Proprietary Novel Drugs businesses. With a network of 45 radiopharmacies in the USA, the Radiopharma business is engaged in manufacturing and supply of radiopharmaceutical products and services. The Allergy Immunotherapy business is involved in the manufacturing and supply of allergic extracts and venom products in the USA and other markets such as Canada, Europe and Australia. Contract Development and Manufacturing of sterile injectables, with facilities in Spokane, USA, and Montreal, Canada, delivers end-to-end manufacturing solutions, including sterile fill-and-finish injectables (liquid and lyophilized), comprehensive ophthalmic products (liquids, ointments and creams) and ampoules. Contract Research Development and Manufacturing business provides end-to-end drug discovery and development services to the pharmaceutical and biotech industries through three world class research centres (two in India and one in France) and a US FDA approved, Active Pharmaceutical Ingredients manufacturing facility in Nanjangud, Karnataka. The Generics business focuses on development, manufacturing and distribution of Solid Dosage Formulations through multiple manufacturing facilities that cater to all the regulated market including USA, Europe and other geographies. Proprietary Novel Drugs is an innovative biopharmaceutical business developing breakthrough therapies in the area of oncology and auto-immune disorders. Jubilant Pharmova 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 visit: www.jubilantpharmova.com.

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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.

Earnings Presentation Q1'FY27

Disclaimer

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Jubilant Bhartia Group has created value across multiple sectors



Strong presence in diverse sectors

- Pharmaceuticals
- Life Science Ingredients
- Performance Polymers
- Food Service (QSR)
- Beverages
- 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

56,000 people across the globe with ~2,200 in North America

Jubilant Pharmova, a diversified pharmaceutical company

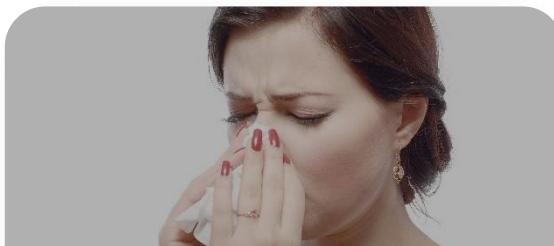


Radiopharma

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

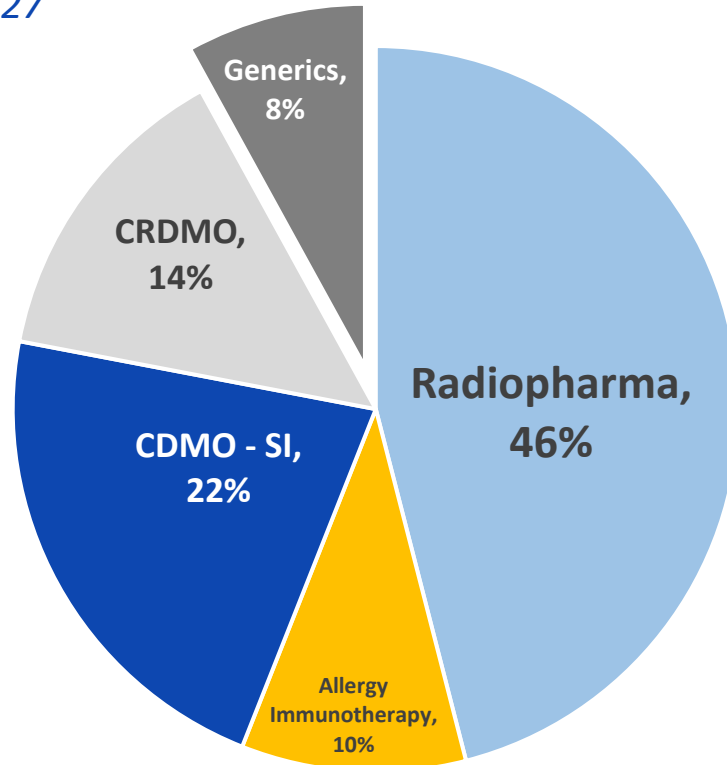
in clinical trials
Developing high potential precision
medicines in Oncology

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

Focus on specialty products & services and Dollar revenues

Business wise Revenue Split

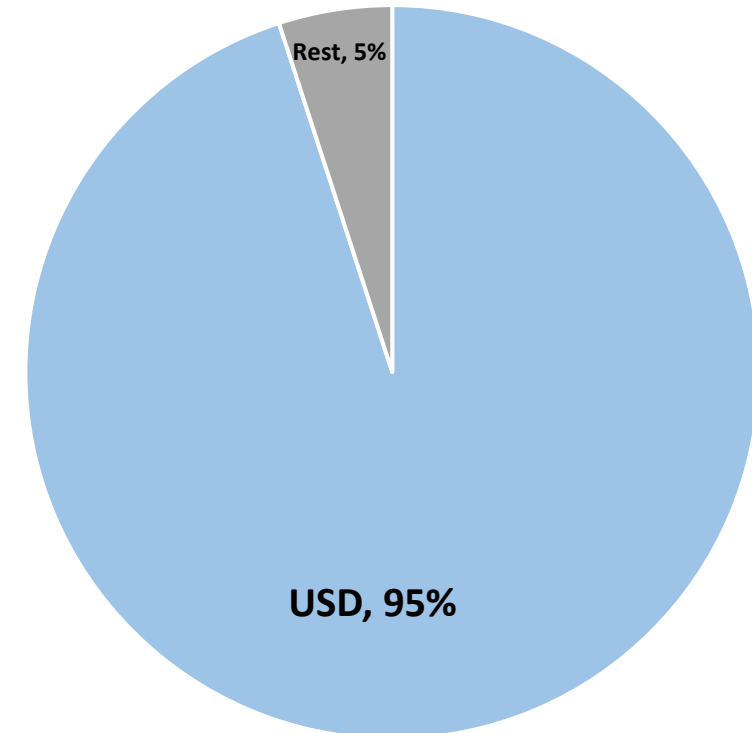
Q1'FY27



**Specialty Products (Radiopharma, Allergy Immunotherapy) – 56%
and Specialty Services (CDMO & CRDMO) – 36%
contribute majority of revenues**

Currency wise Revenue Split

Q1'FY27

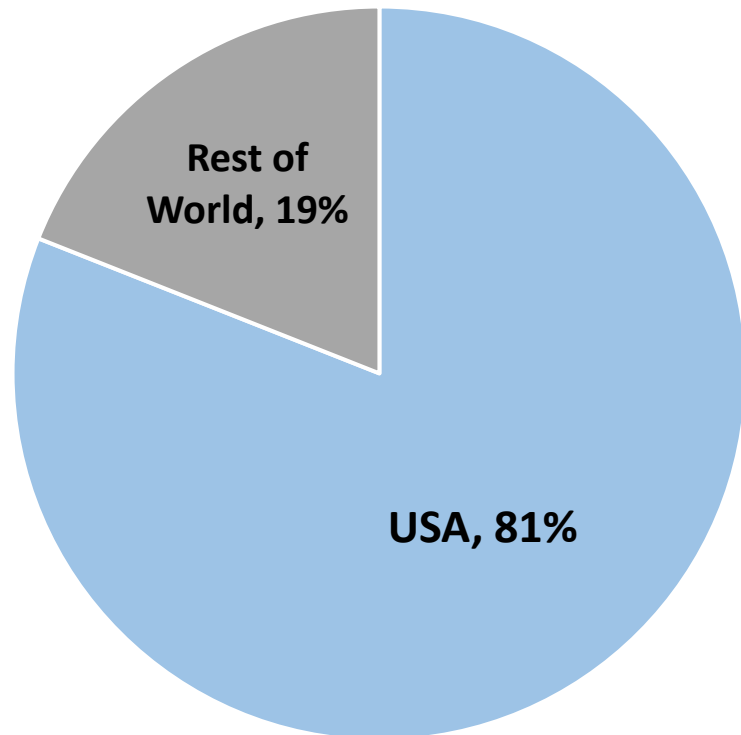


**Majority revenues are
USD denominated**

Minimal risk from US Tariffs

Geography wise Revenue Split

Q1'FY27



US market constitutes majority of revenues

Origin of Goods & Services sold in the US

Q1'FY27



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

* Goods and Services from Canada 14% : Goods 14%, Services 0%

* Goods and Services from India 8% : Goods 3%, Services 5%

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

NORTH AMERICA

Kirkland, Montreal, Canada
CDMO – Sterile Injectables Radiopharmaceuticals



Spokane, Washington, US
CDMO – Sterile Injectables Allergy Immunotherapy



INDIA & EUROPE

Roorkee, Uttarakhand, India - Generics



Nanjangud, Karnataka, India - CDMO API



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 TTM Q1'FY27
2x Revenue	Rs. 6,703 Cr.		Rs. 13,500 Cr.	Rs. 8,608 Cr.
25% EBITDA Margin	~ 15 %		23% to 25%	15%
Zero Net Debt	Rs. 2,457 Cr.		Zero	Rs. 2,332 Cr. End of Q1'FY27
High Teens RoCE	High Single digit		High Teens	11%*

• (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	Progress in Q1'FY27
Radiopharma	Leadership in Ruby-Fill® Launch New PET, SPECT and Therapeutic products (MIBG) Invest in 6 high margin PET Manufacturing facilities in US	Ruby-Fill® growth at 26% Development Progressing Investment on track
Allergy immunotherapy	Strengthen competitive position , develop new products	Gaining Market Share in US
CDMO - Sterile Injectables	Double capacity in Spokane, US	Line 3 Peak Revenue by FY28 Line 4 installation complete
CRDMO	Add large pharma customers Grow CDMO and custom manufacturing in API	Custom manufacturing revenue increasing
Generics	Launch new products in the US and Grow profitable Non-US international business	Launched 2 new products



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

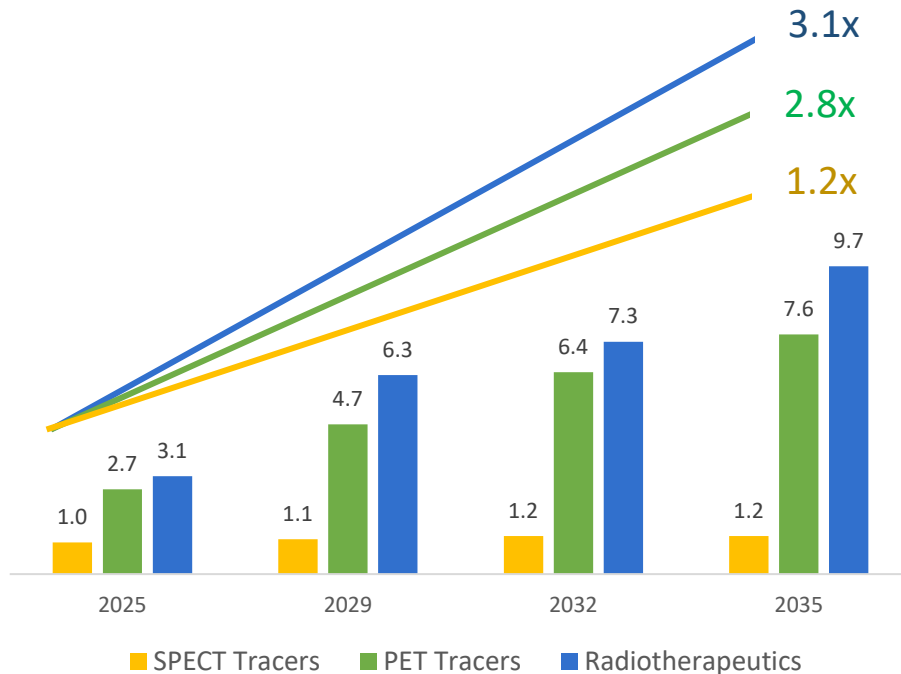
Ruby-Fill[®], Pylarify, Illuccix,
Neuraceq, FDG

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 rapid pace

US Radiopharmaceutical Market USD Bn.



Launch of
advanced
Therapies

New
products in
PET Imaging

Multiple
billion-dollar
M&A deals

Growth Drivers & Trends

- PSMA Therapeutic, Pluvicto for Prostate Cancer ~USD 2.0 Bn.
- PSMA Diagnostics for Prostate Cancer ~ USD 1.8 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 4.1 Bn.)
- BMS and Philochem (USD 1.4 Bn)

PET imaging & advance therapies are driving the market growth

Jubilant Radiopharma – Integrated player with Radiopharma Development, GMP Manufacturing and distribution network

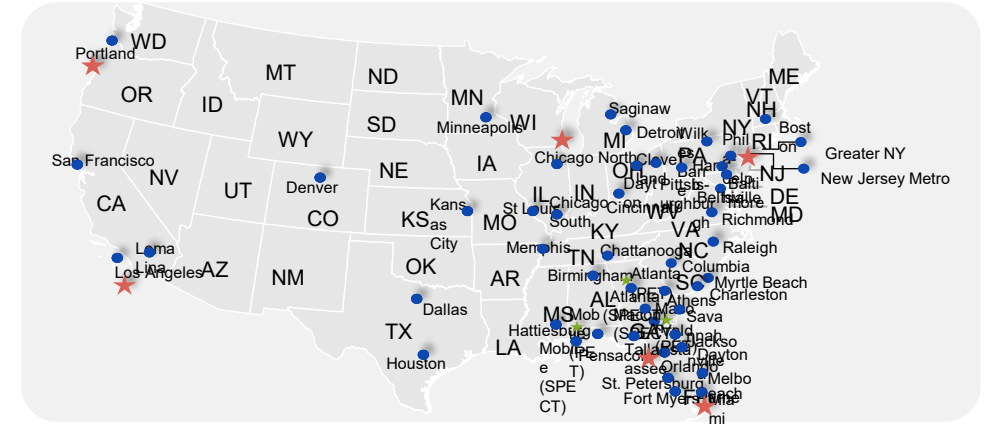


Radiopharmaceutical Plant, Montreal, Canada



- Integrated development and manufacturing site with decades of experience in radiopharmaceutical manufacturing
- Ability to handle multiple isotopes:
 - Diagnostic isotopes : F18, Ga68, Tc99, Rb82, I131Dx
 - Therapeutics isotopes : I131
 - Leader in SPECT products - MAA, DTPA, Cardiac PET product - Ruby-Fill® & Therapeutic I-131HICON®
- Deep pipeline in SPECT / PET assets with upcoming launches
- Final stages of filing for I-131 MIBG (Neuroblastoma)

2nd largest Radiopharmacy Network, US



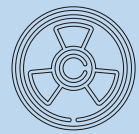
- Network of 42 Nuclear Radiopharmacies and 3 PET Manufacturing facilities
- Serves 1,800 hospitals with 6 customized doses delivered every minute
- Distributes diagnostic radiopharmaceuticals spanning Tc-99, Ga-68, F18, Cu64, I131 and I123
- Expanding the PET manufacturing network from 3 to 9 locations in the US by FY28
- One of the only two pharmacy networks dispensing Lu-177 therapy in the US for Novartis

Solid foundation in SPECT Imaging, Poised to develop PET imaging agents & the supply chain for PET and Therapeutics

Integrated platform across the value chain



Product



Radio -
labeling



Last Mile
Delivery

SPECT

- **Strong and growing** product portfolio

- **2nd largest network of 42 radio pharmacies** distributing ours & industry products across 21 states

- Continuing to build to serve nationally

- Owned fleet of ~400 vehicles to ensure high quality timely delivery and waste management

- **Currently servicing 1,800 customers** at nearly 3,000 individual locations

PET

- **Rapidly growing Cardiac Imaging franchise** and a robust pipeline including pet assets

- **3 operational PET manufacturing facilities with 6 more** in deployment

- Deep experience including ownership and optimization of Sofie platform

- **Integrated with SPECT platform** to ensure a single seamless delivery to multi modality sites and drive operating efficiencies

Therapeutics

- **80% share in I-131**
- Undergoing Clinical Trials for I-131 MIBG in Neuroblastoma

- **Operate one of the few** approved CFR 211 radiopharmaceutical manufacturing sites in North America

- **One of the two networks dispensing Lu177** PSMA Therapy
- Plan to equip more pharmacies in the network to increase reach in Beta emitter therapy dispensing & hub and spoke model for Alphas

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-Gluceptate
		Coronary Artery Disease	Tc99m-Sestamibi
SPECT Dx	Gastrointestinal	Intra-abdominal Infection	Tc99m-Exametazime
	Lung	Pulmonary Embolism	Tc99m-DTPA
		Pulmonary Perfusion	Tc99m-MAA
	Musculoskeletal	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



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

Draximage® DTPA



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

Ruby-Fill®



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® Sodium Iodine I 131 Solution USP



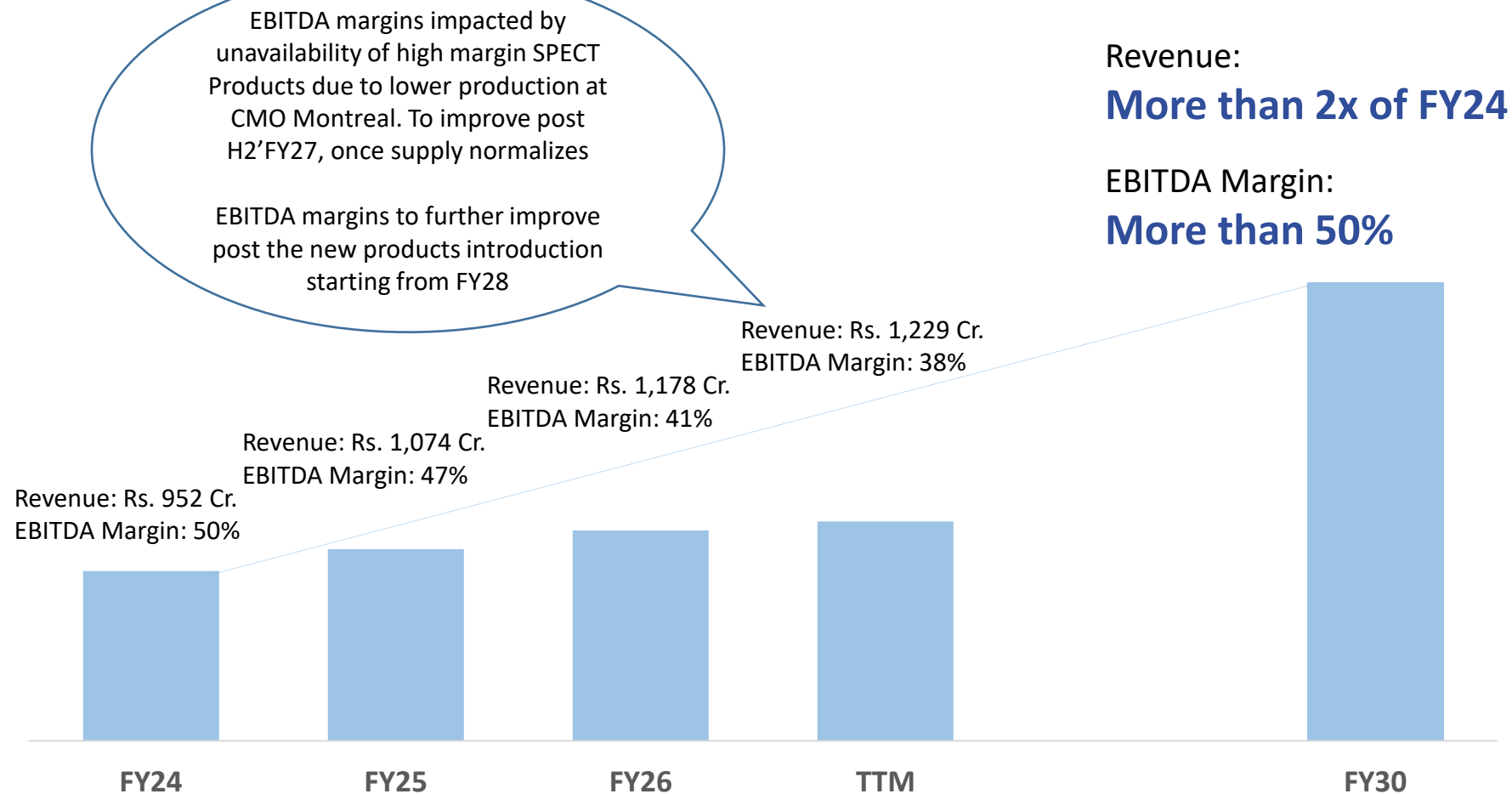
HICON® is a **radioactive therapeutic agent** indicated for the treatment of hyperthyroidism and selected cases of carcinoma of the thyroid. JDI has no direct competition in the US market

Radiopharmaceuticals Financials : Q1'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	271	319	322	19%
EBITDA	126	106	110	(13%)
EBITDA Margin (%)	46%	33%	34%	(1,250) bps

- Revenue grew strongly on back of growth in Ruby-Fill ®
- EBITDA margins decreased YoY due to unavailability of high margins SPECT products
- At CMO Montreal, Commercial batch production started, Batches to be released in Q2'FY27
- Revenue and EBITDA to normalize from H2'FY27 onwards

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®

Growth drivers:

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

Ruby-Fill® Rubidium 82 generator and Elusion System



Competitive advantage

- Longer life per generator (7 weeks vs 6 weeks for peer)
- Better image quality and consistency
- Constant Activity
- Higher number of scans per day vs Fluorine 18 labelled agents
- No additional shielding capex vs Fluorine 18 labelled agents

Current Position

- US Market Size ~ USD 200 + Mn. and growing at 12%
- Market share ~ 25 to 30% and growing
- Margin improving every year

Product Innovation

- AI enabled 3D cardiac blood flow quantification

26% annualized growth in install base in Q1'FY27

Launch new PET and SPECT imaging products with a TAM of USD 535 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	Milestone				
					Products	CMO Selection	Exhibit Batches	End of stability	Launch
	FY28	260	85	3	P1		● →		
					P2	● →			
					P3	● →			
	FY29	275	55	4	P4	● →			
					P5	● →			
					P6	● →			
					P7	● →			
	Total	535	140	7					

Revenue Potential increased to 140 Mn., Progress on track

Launch MIBG by CY27

Growth drivers:

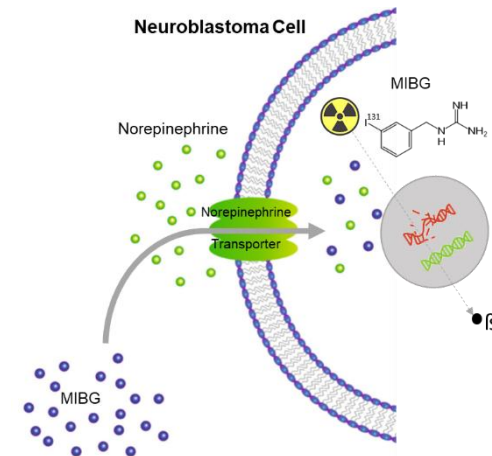
- Ruby-Fill®
- New PET & SPECT products
- MIBG

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.
- NDA filing post FDA meeting by H2'FY27

Radiopharmacy



Radiopharmacies are critical in generating value

SPECT Radiopharmacy



PET Manufacturing Facility



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 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 Manufacturing Facility	# 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

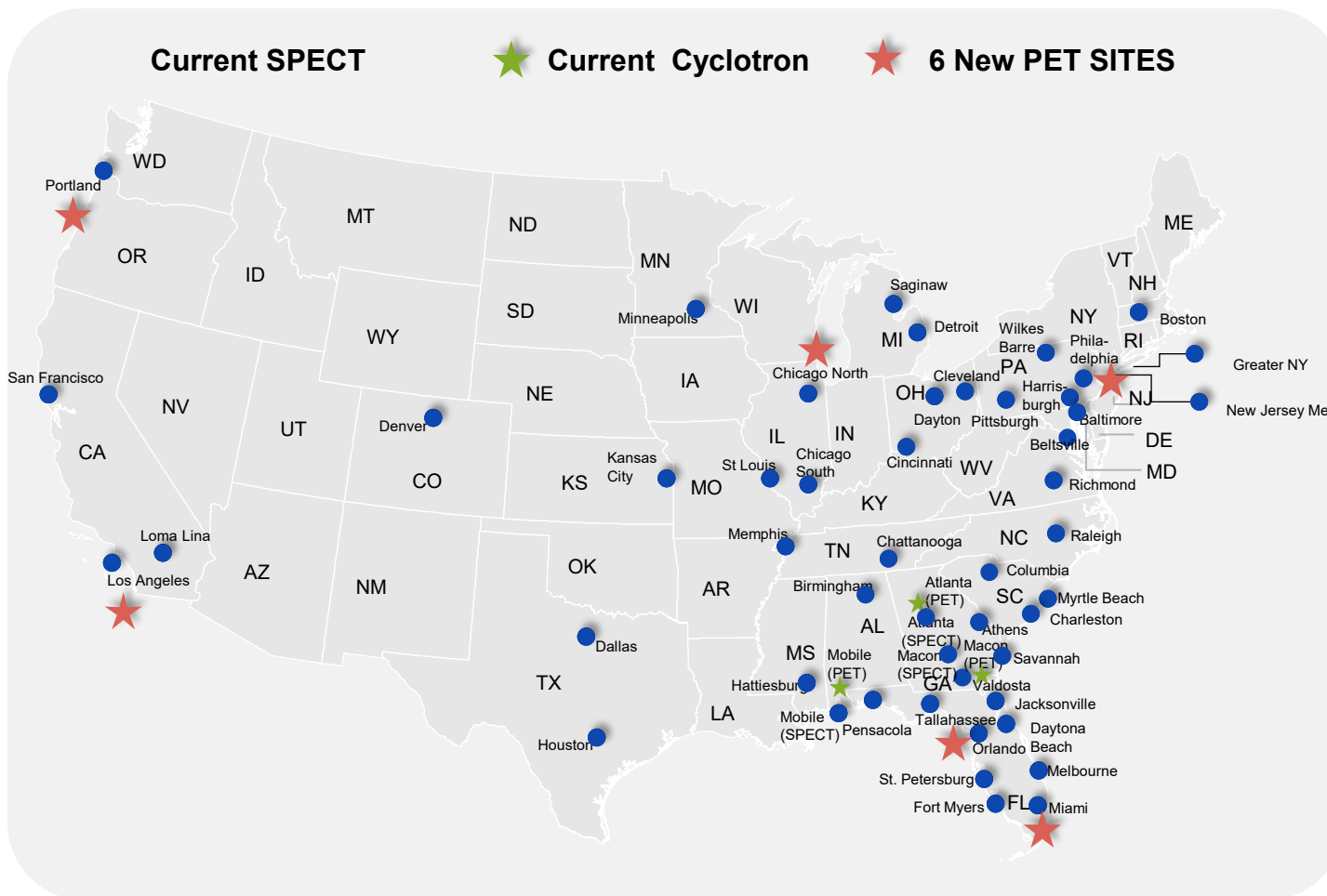


6 customized
doses delivered
every
minute



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

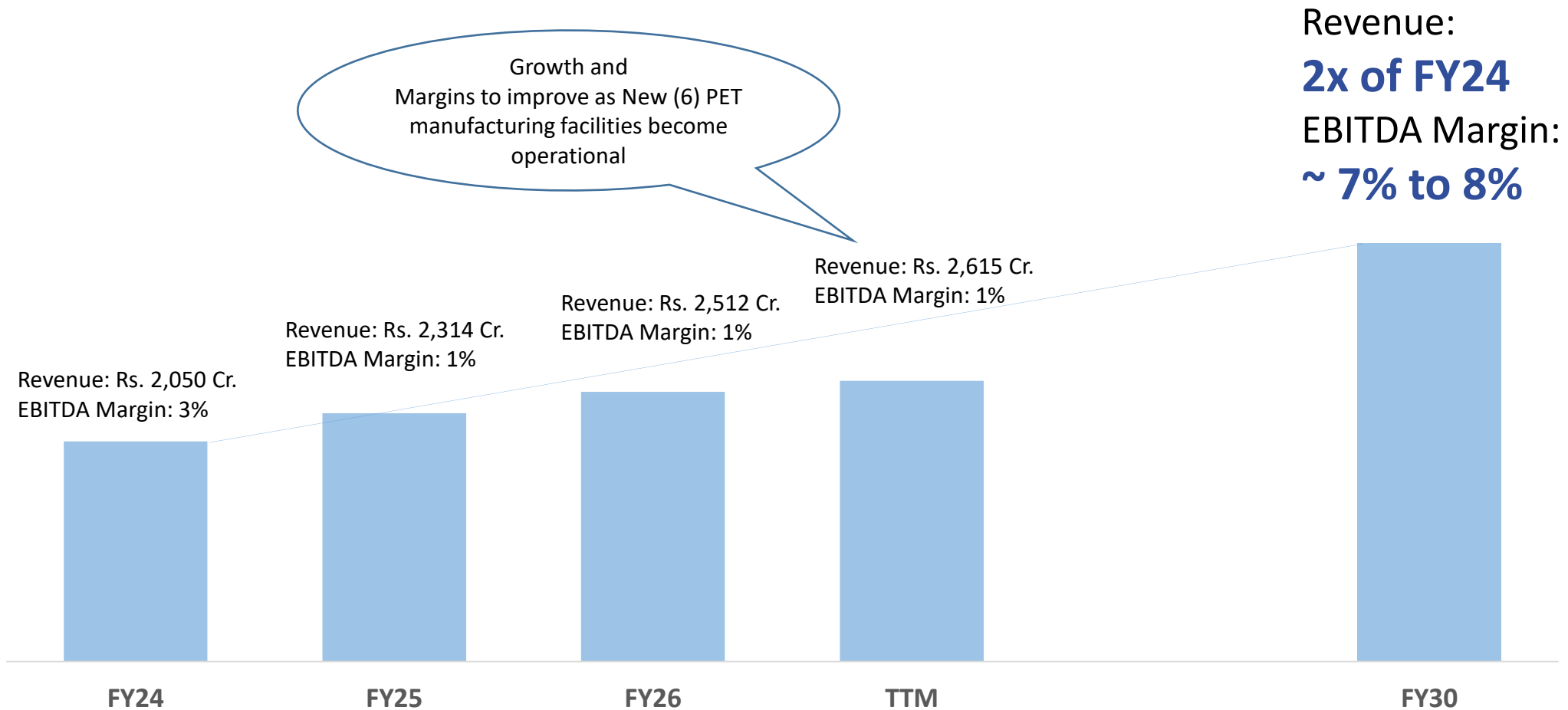
distribution is preferred
from radiopharmacies

Radiopharmacy Financials : Q1'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	598	671	700	17%
EBITDA	10	11	12	19%
EBITDA Margin (%)	2%	2%	2%	0 bps

- Revenue grew YoY on the back of increase in volume from certain PET products. High competitive intensity in SPECT continues
- EBITDA grew YoY on the back of increase in revenue. Expect margin pressure going forward due to high competitive intensity in SPECT products

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



Expanding PET Manufacturing Facilities 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
- **Investment of USD 60 + Mn.**

Continue to witness increase in revenues from the 3 existing PET Manufacturing facilities

A close-up photograph of two bees on a purple flower. The bees are black with yellow stripes. One bee is positioned above the other, both facing towards the left. The flower has many small, delicate purple petals. The background is a soft, out-of-focus green. A semi-transparent dark grey rounded rectangle is centered over the image, containing the text "Allergy Immunotherapy" in white.

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

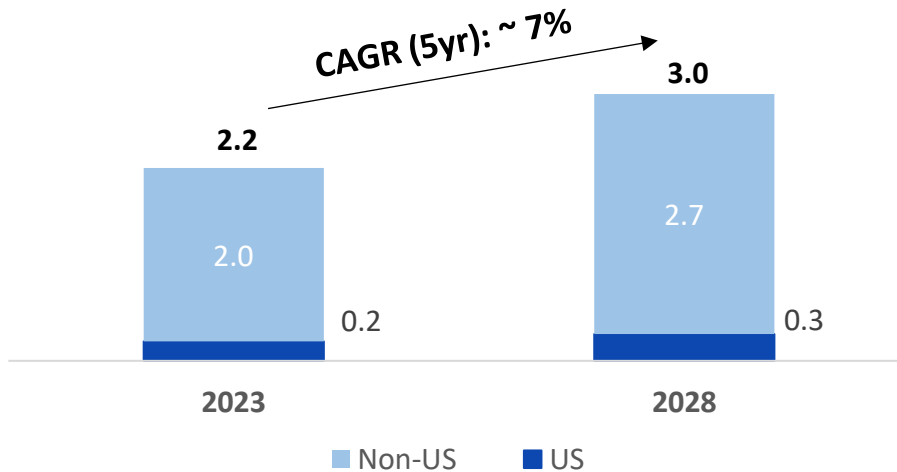


Treatment



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



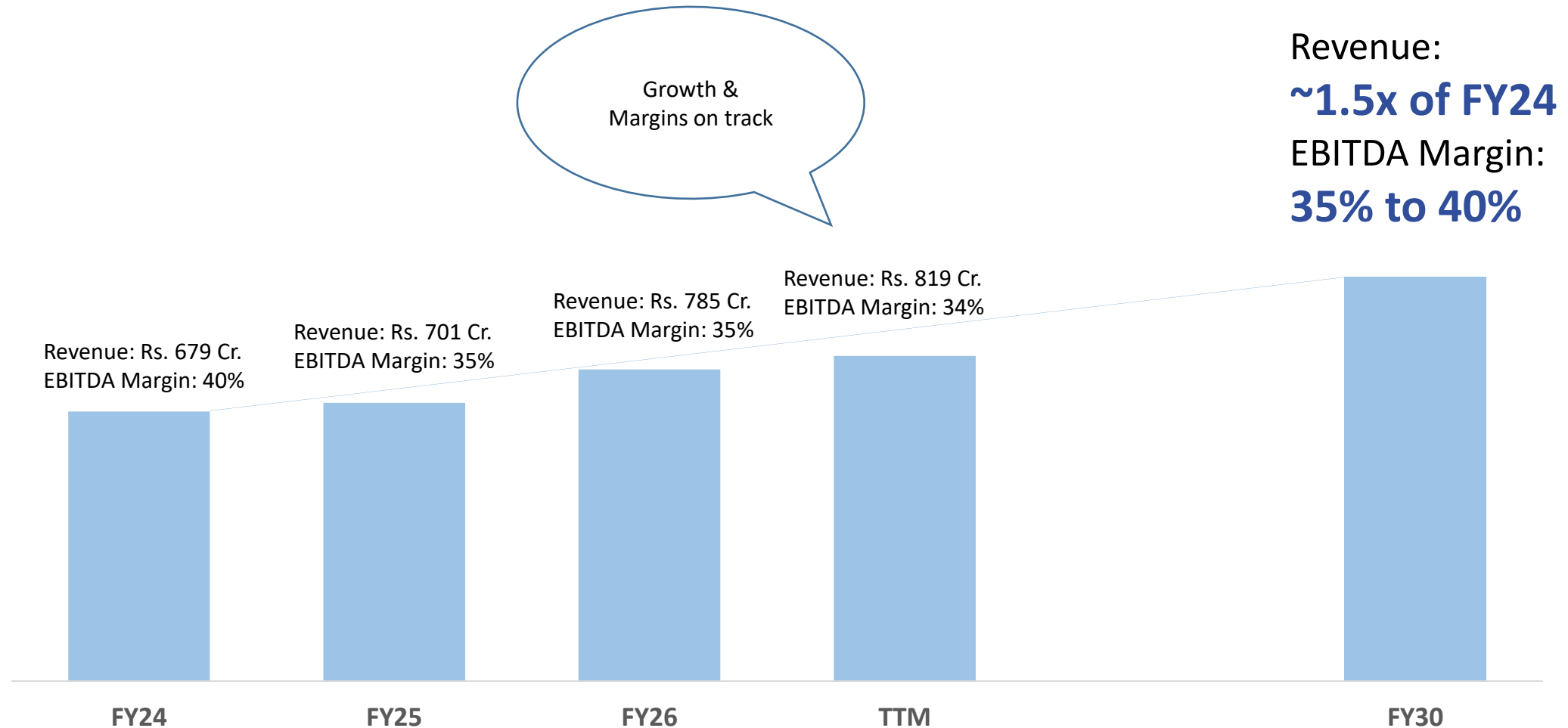
Multiple skin testing systems

Allergy Immunotherapy Financials : Q1'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	181	218	214	18%
EBITDA	63	90	66	5%
EBITDA Margin (%)	35%	41%	31%	(400) bps

- Revenue grew on the back of growth across US & Outside US markets
- EBITDA margin lower YoY due to lower production.

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

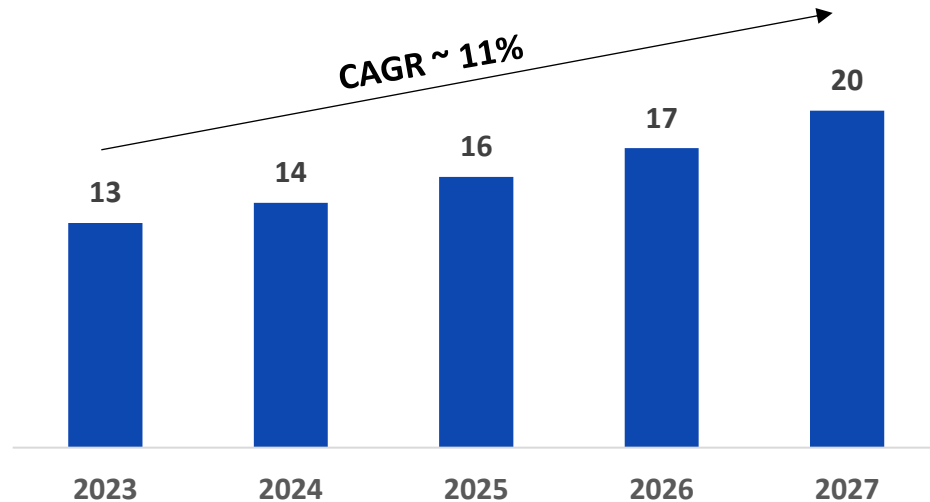
A photograph of a pharmaceutical worker in a cleanroom. The worker is wearing a white full-body protective suit, a hood, and yellow gloves. They are holding a small vial or syringe. The background shows a complex industrial environment with stainless steel equipment, glass partitions, and various pipes and valves. The lighting is bright and even. A semi-transparent dark blue rounded rectangle is overlaid on the center of the image, containing the text "CDMO - Sterile Injectables" in white.

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 and Ointments) 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.

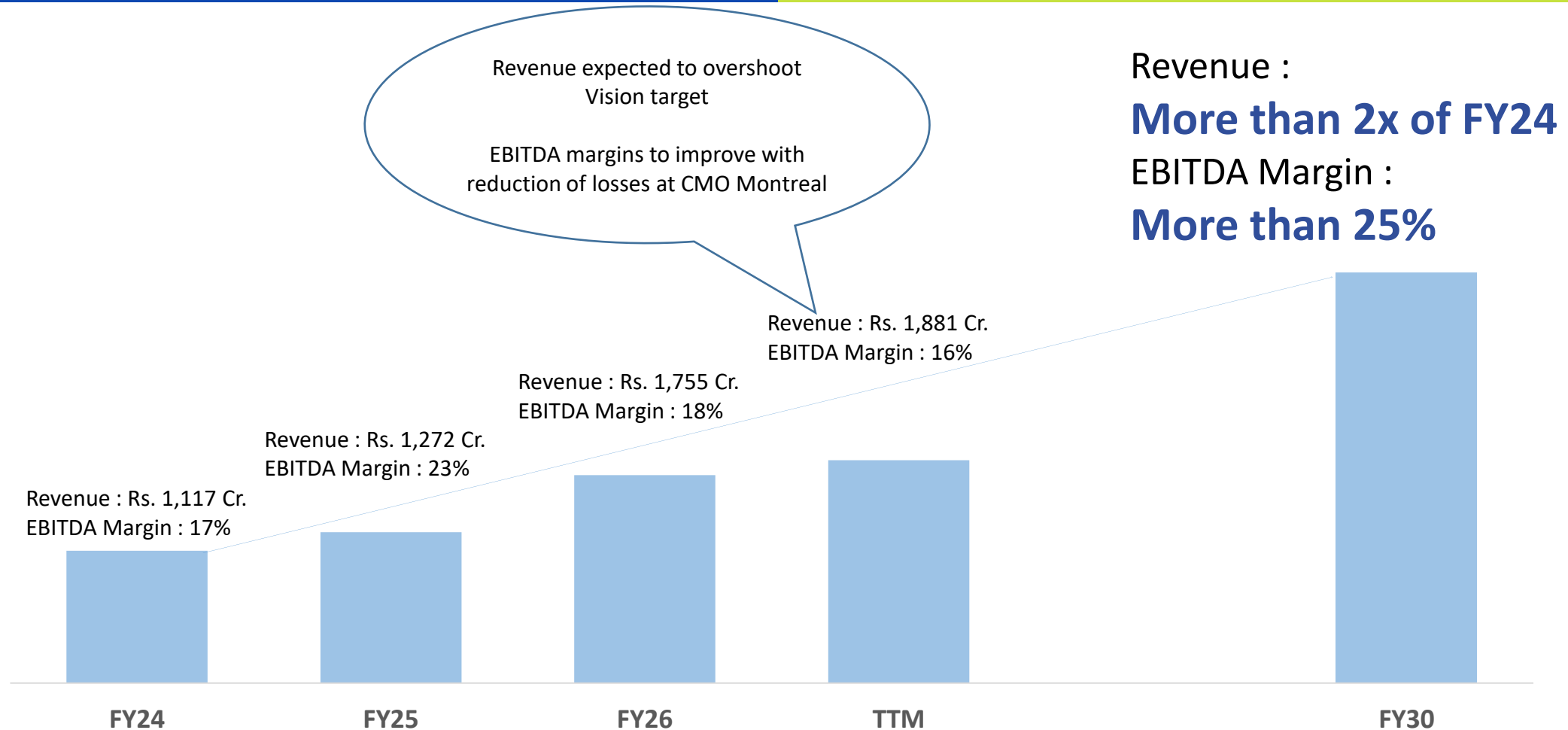
CDMO Sterile Injectables Financials : Q1'FY27

TOTAL Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	370	535	496	34%
EBITDA	62	91	45	(28%)
EBITDA Margin (%)	17%	17%	9%	(780) bps

SPOKANE Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	346	532	494	43%
EBITDA	78	166	117	49%
EBITDA Margin (%)	23%	31%	24%	100 bps

- Revenue grew strongly on YoY due to incremental revenues from Line 3 at Spokane. Revenue lower QoQ due to planned maintenance shutdown
- EBITDA margin lower YoY due to negligible third-party revenues & higher operating expenses including incremental remediation cost at CMO Montreal
- Spokane revenue grew strongly on the back of scale up of Line 3, scaling up faster than the industry benchmark. Revenue lower QoQ due to planned maintenance shutdown
- Spokane EBITDA margin expanded YoY on the back of best-in-class revenue growth

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

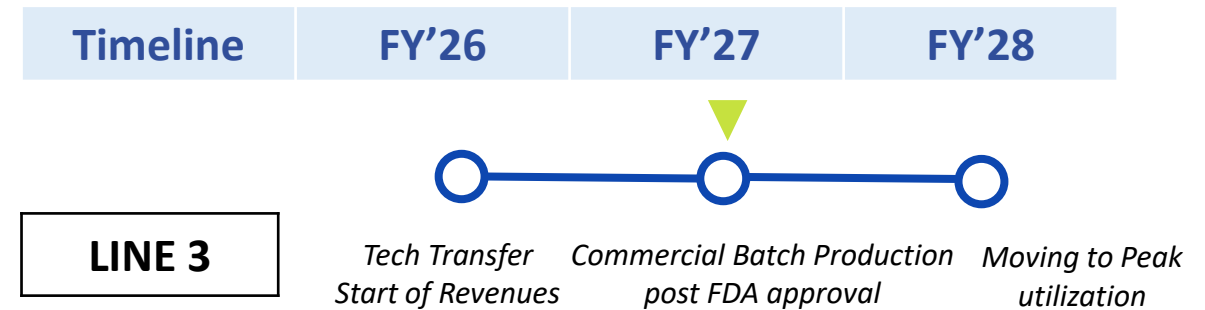


Line 3 Technology transfer revenues continue to grow

Commercial Batch Production expected to start in FY27

Growth driver:

- Doubling Capacity at Spokane



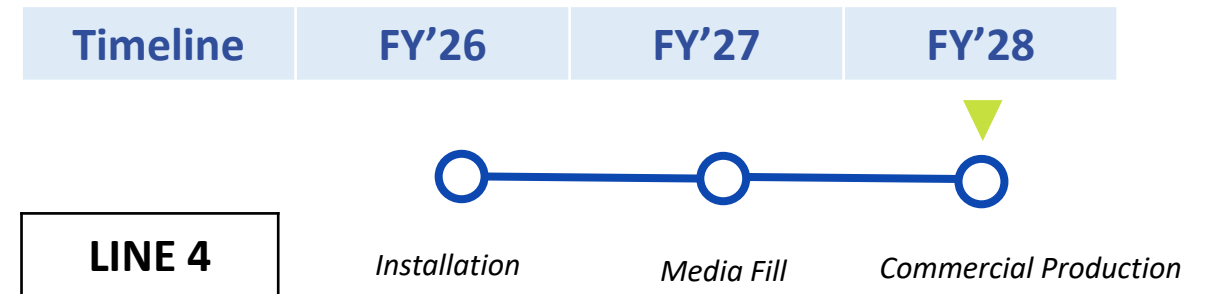
- Line 3 revenues scale up is one of the fastest in the industry on the back of 10+ technology transfer programs
- Expect commercial batch production to start in FY27; To reach full utilization in 3 ½ years
- Peak revenue potential of USD 80 to 90 Mn.

Line 4 installation on track

Technology transfer revenues expected to start in FY27

Growth driver:

- Doubling Capacity at Spokane



- Line 4 installation complete
- Building a strong order book pipeline
- Expect technology transfer revenues to start in Q4'FY27
- To reach full utilization in 3 ½ years
- Peak revenue potential of USD 80 to 90 Mn.

Line 5 to generate tech transfer revenues from FY29

Growth driver:

- New Isolator Line at Montreal



New Isolator Fill & Finish Line (Line 5)

- Construction started; Orders placed for Plant & Machinery
- Capex at USD 114 Mn., Concessional loan at USD 35 Mn.
- Expect Technology transfer revenue to start in FY29
- Scale up of revenue will be similar to Line 3 & 4

Existing Line

- Production started for SPECT Radiopharmaceutical products.
- Remediation work is in progress, Engaged with FDA
- Continue lean operations in FY27 & FY28, till Line 5 becomes operational in FY29

With Line 3,4&5, We are becoming a specialized CDMO delivering complex biologics formulations for Innovators



Traditional CDMO

- **Product Mix** ~ 80% small molecules & **20% Biologics** with low formulation complexity
- **Capability** viewed as **volume driven CDMO**
- **Quality of Revenue: Lower technology transfer revenue.** Programs are easier to onboard and quicker to qualify

Line 3, 4 & 5

- **Product Mix** projected at 20% small molecules & **80% Biologics** with complex formulations (e.g. advanced processing requirements & Technology Transfer packages, high-complexity and high value APIs)
- **Capability** includes **specialised filling, tighter aseptic processing windows** & stringent environment controls
- **Quality of Revenue: Higher technology transfer revenue** model. Programs are difficult to onboard & longer to qualify, once commercialised, more durable

Onboarded one of the World's Largest Oncology products on Line 3



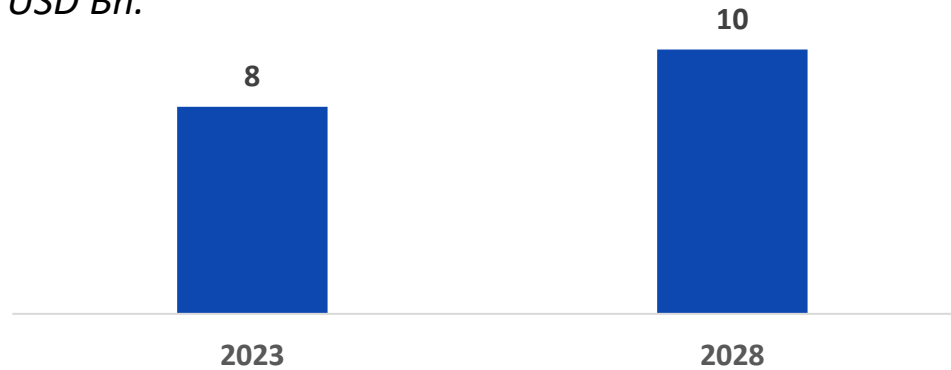
CRDMO: Drug Discovery Services, CDMO API

CRDMO: Drug Discovery, CDMO - API

India uniquely positioned to benefit from Friendshoring

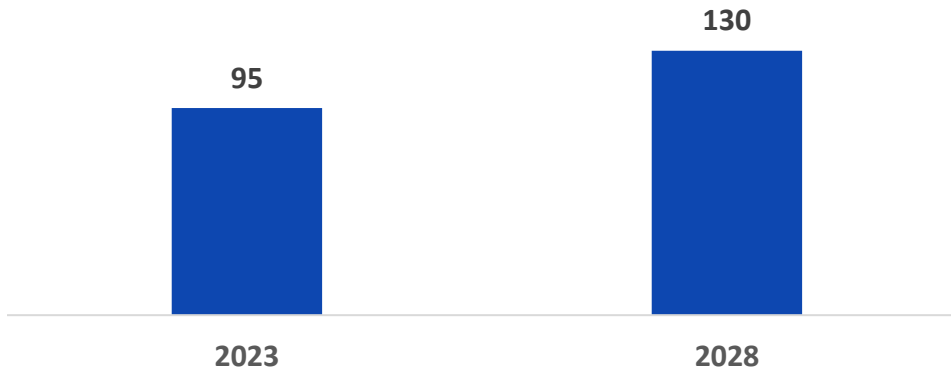
Drug Discovery Services Market Size

USD Bn.



CDMO API Market Size

USD Bn.



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



- ~ **One third revenue contribution** from Large Pharma accounts
- **Indian Leader for “Integrated Drug Discovery”**, with a track record of 100+ 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 during the year

...with Integrated Discovery & Development Infrastructure

Drug Discovery

Integrated Drug Discovery Center (Bengaluru, India)



~400 Scientists
120+ IDD programs

- Target ID to pre-clinical candidate selection with best-in-class drug discovery timelines
- Structural Biology, CADD, In Vitro, In Vivo Biology, DMPK, Toxicology
- AAALAC-accredited vivarium

Chemistry Innovation Research Center (Delhi NCR, India)



~750 Scientists
6 Centers of Excellence

- Synthetic, Medicinal, Analytical & Computational Chemistry; DMPK, Biology
- Expertise in Degraders, Peptides, Lipids, Solid State, Library Synthesis, Photo Redox, Carbohydrates

Center of Excellence for Biologics & ADCs (St. Julien, France)



~35 Scientists
In the Pharma Hub of Europe

- ADCs - Full discovery loop
- Biologics & Immune-Oncology expertise
- Antibody engineering, linker chemistry, payload conjugation

CDMO and API

Advanced Center for PR&D and Scale-Up (Delhi NCR, India)



~100 Scientists
PRD & Non GMP Scale up

- Greater than 1 KL capacity
- 15 reactors (20 to 250 L)
- Multiple PRD programs ranging from Phase 1 to Phase 3
- SS, GLR, Hastelloy Reactors
- Material Generation upto 25KG
- FFS Compatible

GMP CDMO & API Manufacturing Facility (Nanjangud, India)



900+ MT production capacity
785 KL reactor capacity

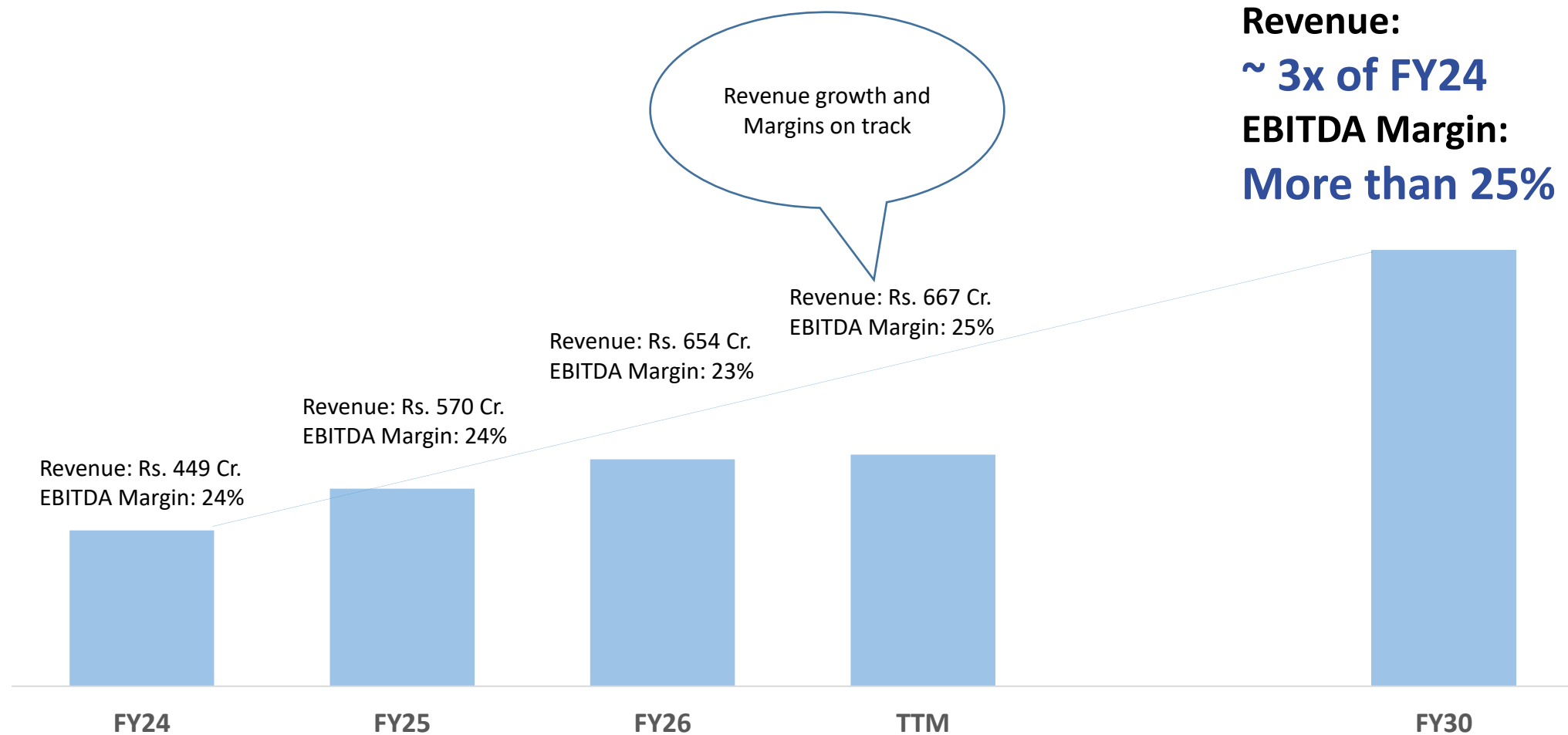
- 6 Plants, Kilo Lab & Pilot Plant
- Potent API expertise
- OEB Class 1-4 API potency
- 50 gm - 1.2 Ton GMP Batch
- Digitally Enabled (DCS + QA/QC)

Drug Discovery Financials : Q1'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	161	162	174	8%
EBITDA	32	42	45	43%
<i>EBITDA Margin (%)</i>	<i>20%</i>	<i>26%</i>	<i>26%</i>	<i>630 bps</i>

- Revenue increased YoY on the back of increase in revenue from biotech and large pharma customers
- EBITDA margins improved YoY due to better revenue mix and forex gain

Drug Discovery Vision 2030 : Triple revenues & maintain profitability



Drug Discovery Services: Leverage Large Pharma potential



Growth driver:

- Add Large Pharma



Biosecure Act

- **Biosecure ACT becomes law** in the Unites states
- Federal agencies must not enter in contract with a biotechnology company of concern

- Execute strategy on Large Pharma
- Build Footprint in EU
- Introduce 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

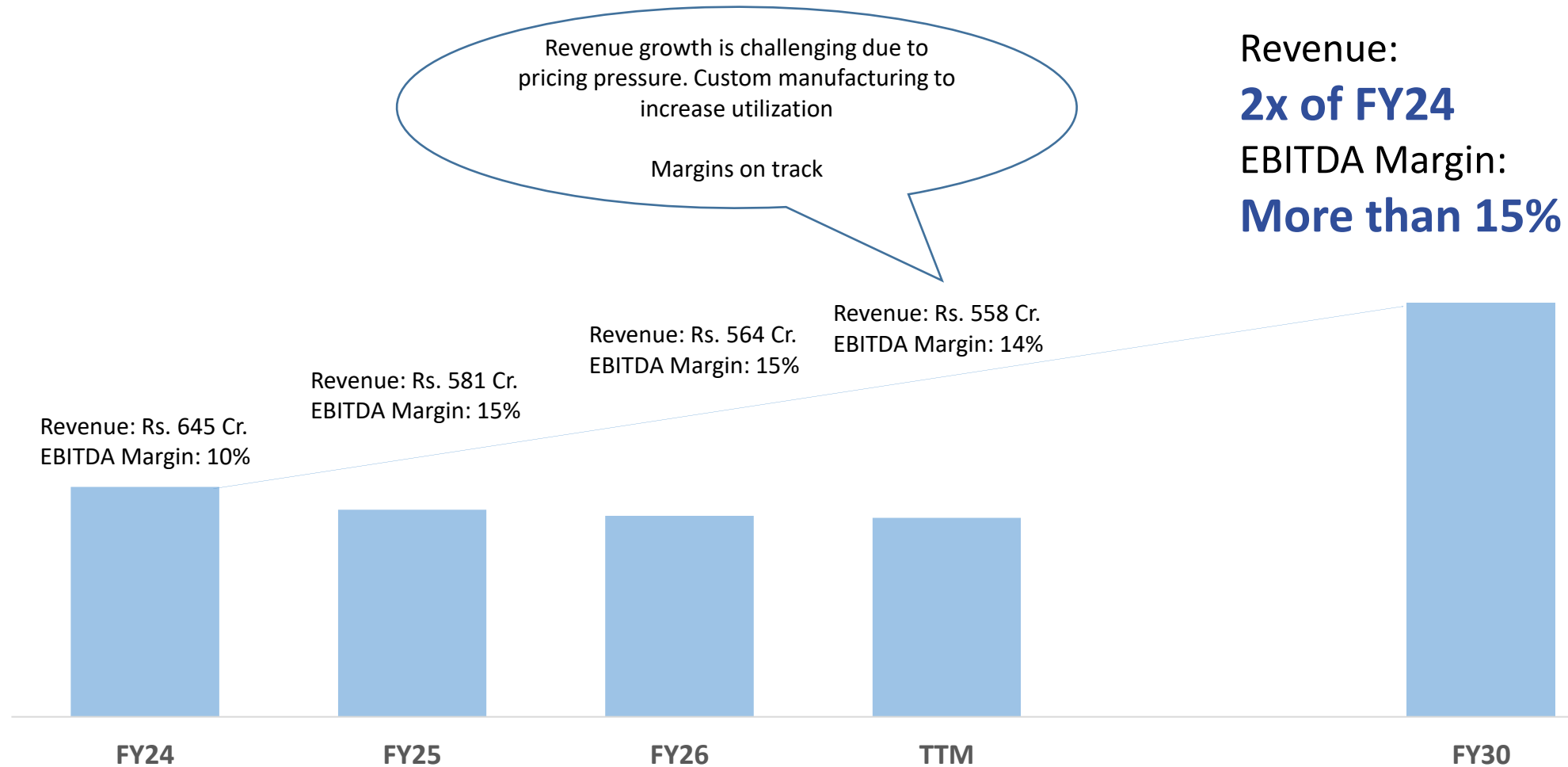
Integration complete; Investing in Business development team

API Financials : Q1'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	141	156	135	(4%)
EBITDA	22	22	19	(12%)
EBITDA Margin (%)	15%	14%	14%	(130) bps

- Industry wide pricing pressure continues. Focus to increase revenue mix from Custom manufacturing
- EBITDA margins marginally lower YoY due to lower revenue and increase in raw material cost
- Expect margin pressure in FY27 due to raw material cost inflation on account of current ongoing middle east conflict. Focusing on high margin products to offset the impact.

API Vision 2030 : Double revenues and increase profitability



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

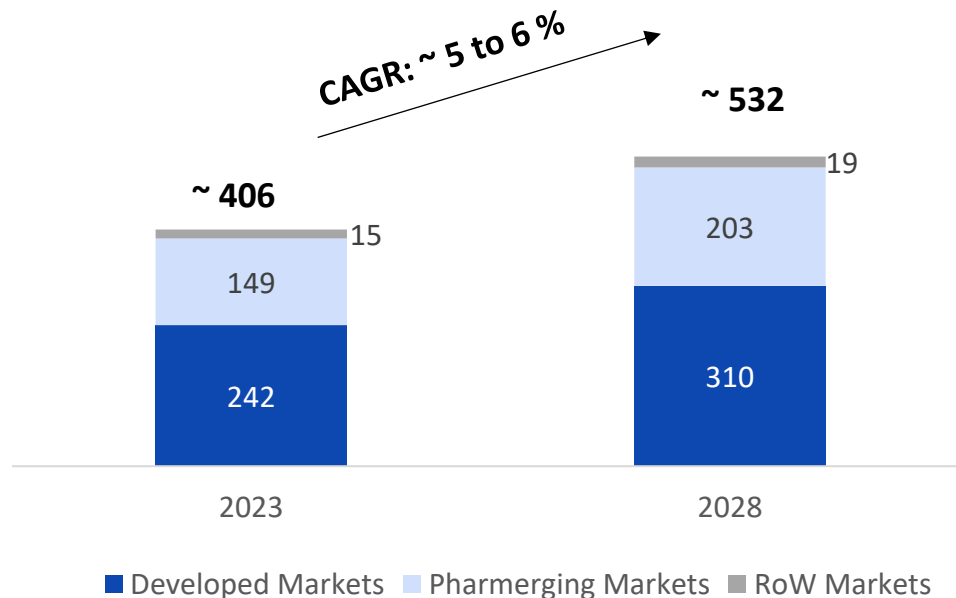
- Completed sale and transfer of API business to “Jubilant Biosys”, wholly owned subsidiary of company
- Combined platform to improve operational efficiency and superior brand recall of “Jubilant Biosys”
- Increase asset utilization of API business by improving revenue mix towards Custom manufacturing & CDMO



Generics

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

Generics Market USD Bn



Growth Drivers and Trends

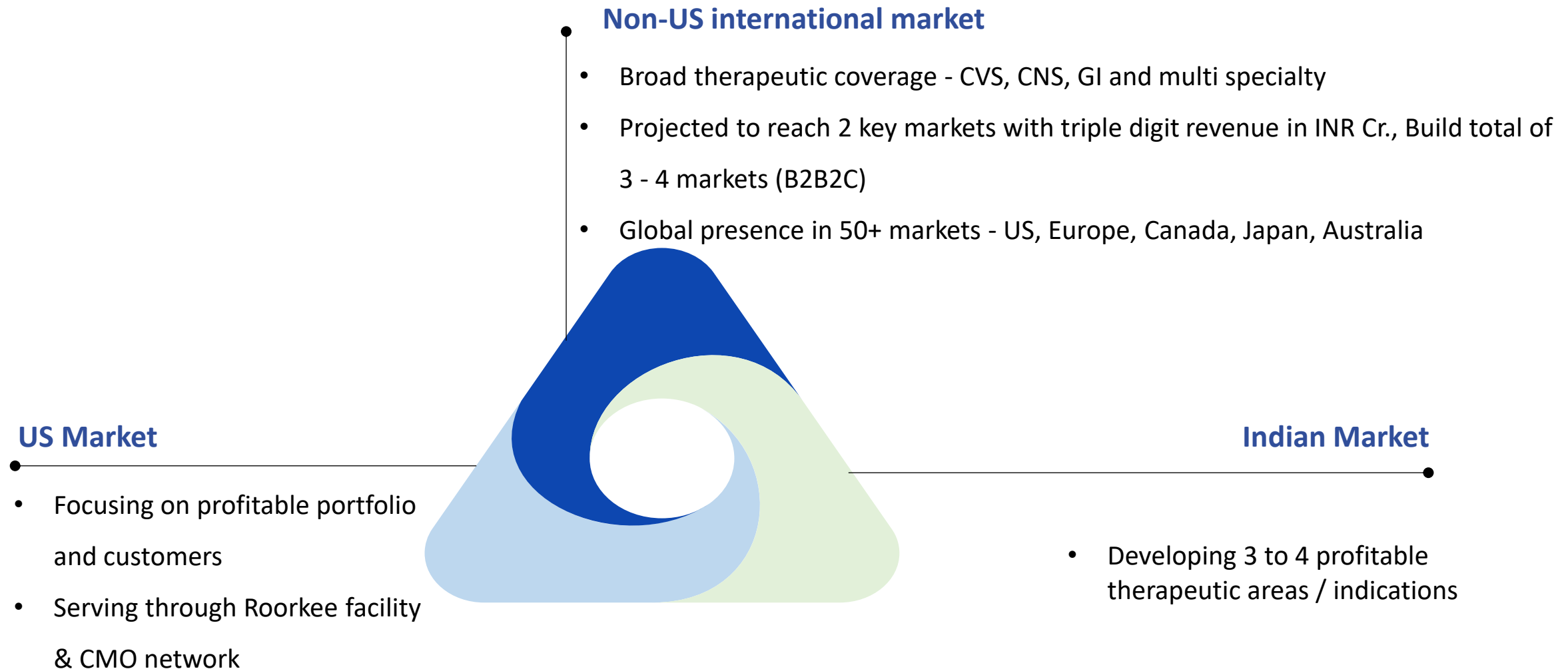
Developed Market

- US market to grow at 2%
- Non-US market to grow by 5 - 7%

India Market

- India market to grow ~ 8-9%
- Brand building and in-clinic effectiveness are key drivers

We are building a growing, profitable & agile business model



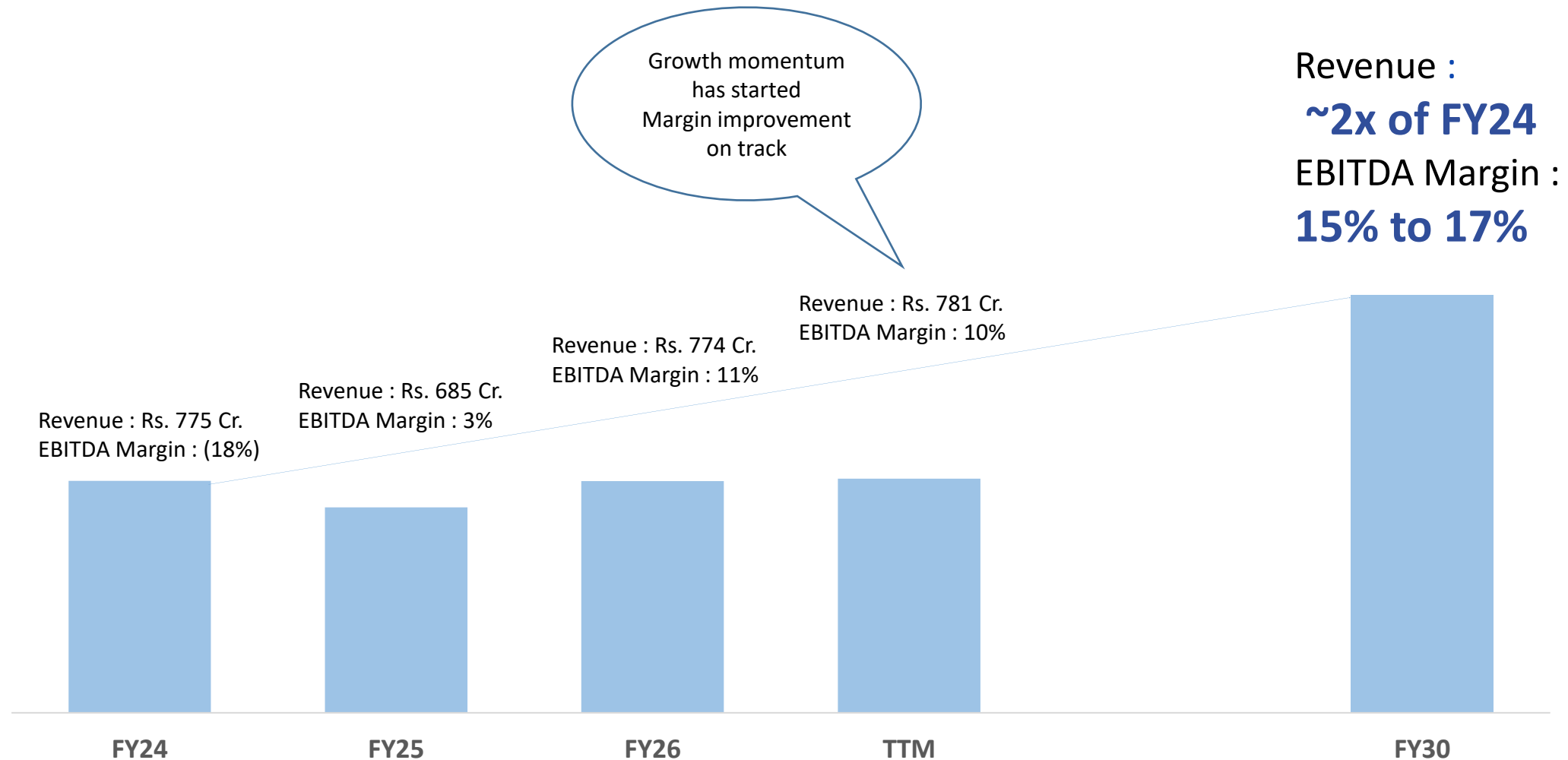
Generics Financials : Q1'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	166	214	173	4%
EBITDA	12	32	4	(65%)
EBITDA Margin (%)	7%	15%	2%	(480) bps

- Revenue grew marginally on YoY. Launched 2 new products in Q1'FY27.
- EBITDA margins lower YoY due to change in product mix. Trailing 12 month margin at ~ 10%. Expect margins to improve in subsequent quarters

Generics Vision 2030:

Reach top quartile profitability for similar size companies



Generics Growth Drivers



Launch 6 to 8 new products annually

- Relaunch dormant ANDAs from Roorkee and CMO network
- Secure ANDAs approvals
- In license and acquire targeted ANDAs



Grow the profitable Non-US international market

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



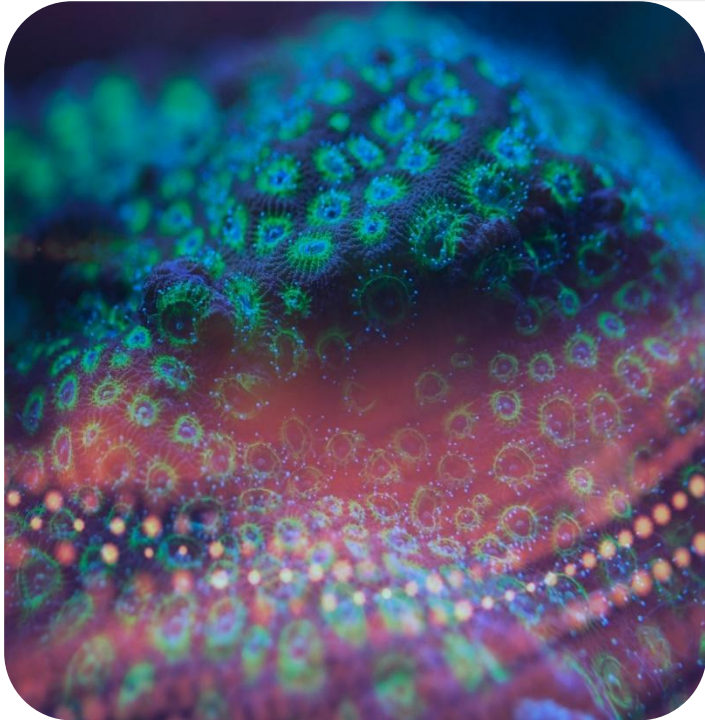
Build branded business

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



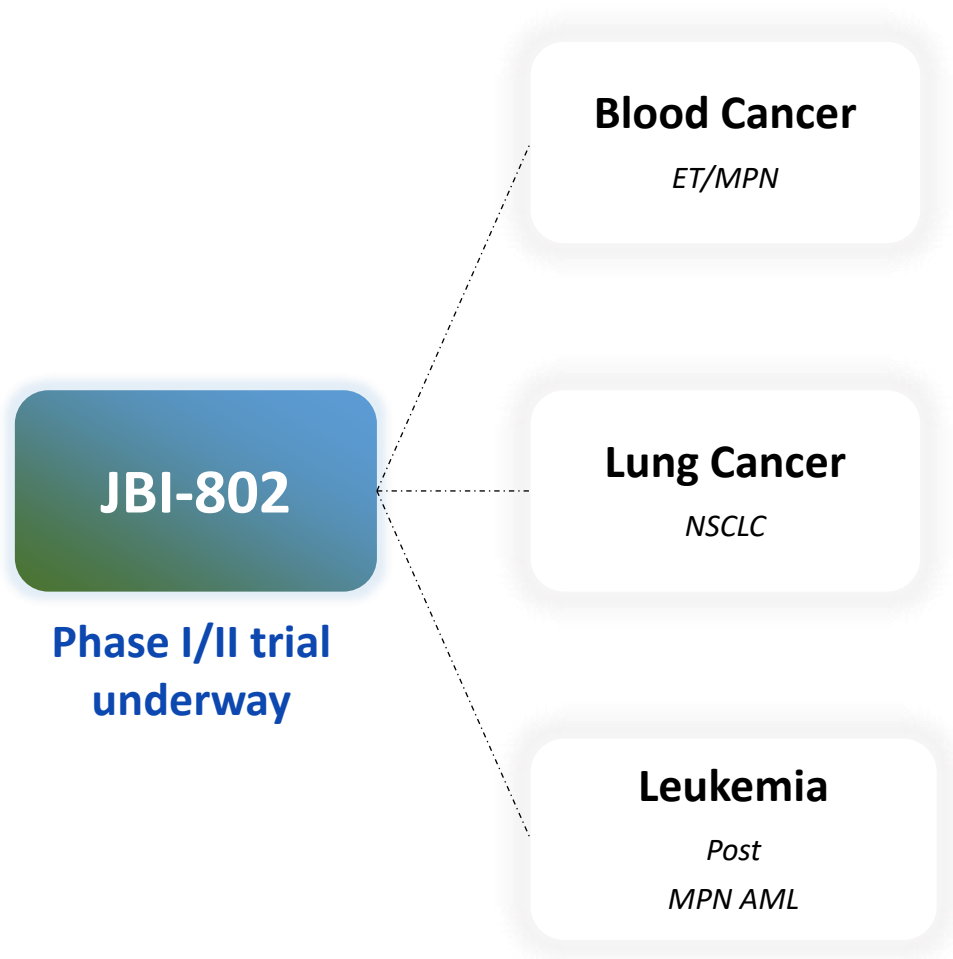
Proprietary Novel Drugs

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 leading oncologists** from World renowned institutions
- **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 I/II trial underway**
 - Highly differentiated for safety and efficacy than peers
 - Total Addressable Market in US ~ USD 5 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.

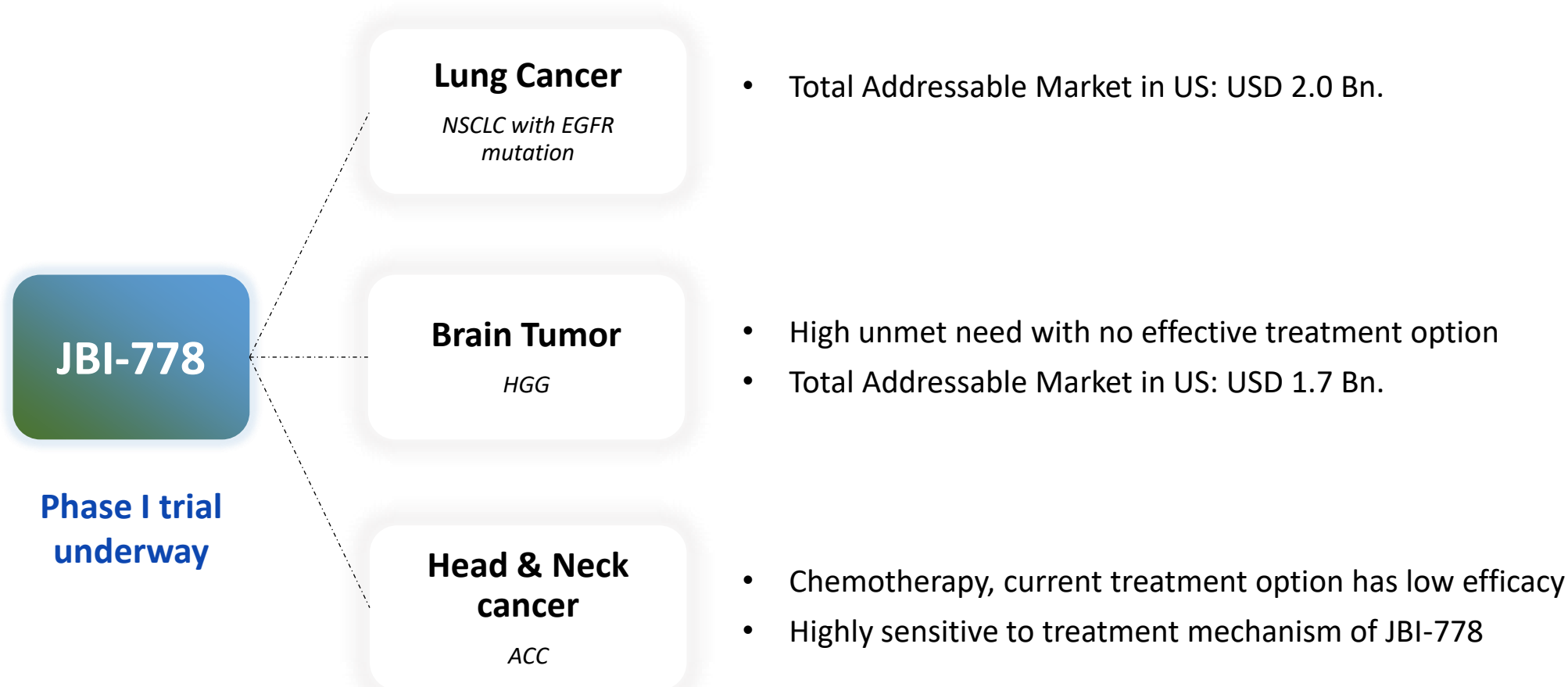
JBI -802 has demonstrated rapid, durable platelet normalization in Essential Thrombocythemia patients



Image only for reference

- **Essential Thrombocythemia (ET)** is a WHO-defined myeloproliferative neoplasm (blood cancer) characterized by elevated platelets, with risks of thrombosis e.g., stroke, bleeding (hemorrhage) and progression to myelofibrosis or Leukemia, representing **a meaningful unmet need and commercial opportunity**
- Anagrelide is the only FDA approved drug for ET and has a boxed warning. Other drugs such as Hydroxyurea are used off-label but have poor patient outcomes
- No FDA approvals for ~30 years which means **a field ripe for targeted therapy with ~150,000 patients in U.S alone**
- **Early Phase I/II data from JBI-802 demonstrates rapid, durable, dose-dependent platelet reduction, with potential superior safety profile**

JB1-778 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'FY27

Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	0	0	0	
EBITDA	(6)	(7)	(8)	(23%)

- 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'FY27

Solid revenue growth (YoY) across all business segments



Particulars (Rs. Cr.)	Q1'FY26	Q4'FY26	Q1'FY27	Y-o-Y
Revenue	1,901	2,290	2,229	17%
Other Income	12	24	20	
Total Income	1,913	2,314	2,249	18%
EBITDA	302	363	268	(11%)
<i>EBITDA Margin (%)</i>	<i>15.8%</i>	<i>15.7%</i>	<i>11.9%</i>	<i>(385) bps</i>
Exceptional Income / (expense)	0	(14)	0	
PBT	154	176	86	
PBT Margin	8.1%	7.6%	3.8%	
Normalised PBT¹	154	190	86	(44%)
Normalised PBT Margin	8.1%	8.2%	3.8%	(423) bps
Reported PAT	103	119	56	(45%)
Reported PAT Margin	5.4%	5.2%	2.5%	(285) Bps
Normalised PAT²	103	129	56	(45%)
Normalised PAT Margin	5.4%	5.6%	2.5%	(285) bps

- **Revenue grew YoY** on the back of strong performance across all business segments, with CDMO Sterile Injectables delivering particularly robust growth
- **Other income for Q1'FY27** includes **grant income** of Rs. **5.6 Cr.** for Line 3, which shall continue for more than 20 years and upto 30 years.
- **EBITDA decreased YoY**, particularly due to unavailability of SPECT products in Radiopharmaceuticals & negligible third-party revenues and higher operating expenses including incremental remediation cost at CMO Montreal
- **Reported PAT decreased YoY** due to lower operating profitability and increase in depreciation for Line 3 in Spokane

1. Normalised PBT is after adjusting for Exceptional items

2. Normalised PAT is after adjusting for Exceptional items and tax

Key Ratios

Net Debt / Ebitda to remain range bound

Particulars (Rs. Cr.)	Mar 31, 2026	June 30, 2026
Net Debt (On constant currency, Net of DIC)	1,952	2,338
Net Debt / Equity	0.28	0.33
Net Debt / EBITDA (TTM)	1.5	1.8
Interest Coverage Ratio	6.3	6.0
Long Term Capex Creditors	711	710

- **Net debt / Ebitda to remain range bound**
- **Key ongoing Capex Projects Include**
 - New (6) PET Manufacturing Facilities
 - CDMO SI, Spokane Line 4
 - CDMO SI, Montreal Line 5
 - Allergy Immunotherapy facility upgrade
 - New Product Development

Sustainability



NSE
Sustainability
Ratings & Analytics
Category I ERP

ESG Score 72 % Leader



dun & bradstreet
ESG Horizons:
Now and Next

Recognized – Leading ESG entity



BRONZE | Top 35%
ecovadis
Sustainability Rating
JUL 2025

EcoVadis Score 61 %



S&P Global
Dow Jones
Sustainability Indexes

DJSI Score 57%



Crisil
ESG Ratings
& Analytics

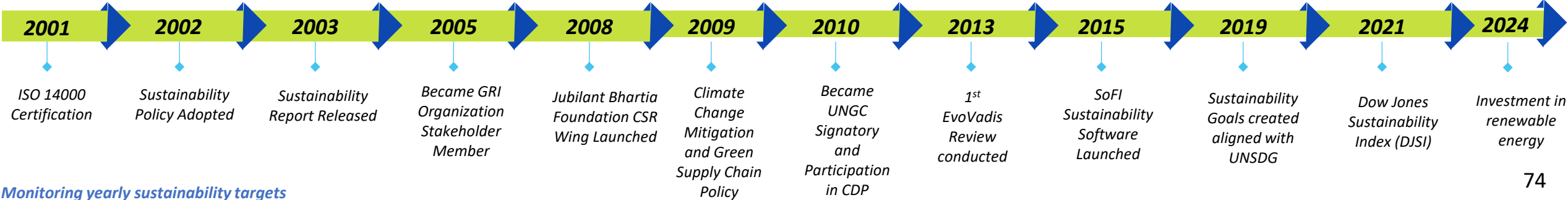
ESG Score 63%

**FY25
Sustainability
Report published**

Assured by EY



**FY25 Sustainability
Linked Loan KPIs
Assurance
completed by EY**



Summary – Q1'FY27

1

Radio Pharmaceuticals : Ruby-Fill® maintaining **growth momentum**. Revenue and EBITDA to normalize from H2'FY27
Radio Pharmacies : Competitive intensity higher in SPECT, **PET products revenue** continue to grow

2

Allergy Immunotherapy : Revenue grew YoY; EBITDA margins decreased due to lower production.

3

CDMO Sterile Injectable : **Strong revenue growth from Line 3 tech transfer programs**.
Commercial batch production started for existing SPECT radiopharmaceutical products at CDMO Montreal

4

CRDMO DDS: Steady growth & profitability improvement amid intensifying competition. **Medium term outlook continues to be positive**
CRDMO API : Focus on profitable products. **Taking initiatives to build custom manufacturing business**

5

Generics : Improved **growth & profitability outlook**

6

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

Financial Results Table

Total Income (Rs. Cr.)	Q1'FY26		Q4'FY26		Q1'FY27	growth
Revenue (A)	1,901		2,290		2,229	17%
a. Radiopharma	869		990		1,022	18%
<i>Radiopharmaceuticals</i>	271		319		322	19%
<i>Radiopharmacies</i>	598		671		700	17%
b. Allergy Immunotherapy	181		218		214	18%
c. CDMO Sterile Injectables	370		535		496	34%
d. CRDMO	302		318		309	2%
<i>Drug Discovery Services</i>	161		162		174	8%
<i>CDMO – API</i>	141		156		135	(4%)
e. Generics	166		214		173	4%
f. Proprietary Novel Drugs	0		0		0	
<i>Unallocable Corporate Income</i>	13		15		14	11%
Other Income (B)	12		24		20	66%
Total Income (A+B)	1,913		2,314		2,249	18%

EBITDA (Rs. Cr.)	Q1'FY26	Margin	Q4'FY26	Margin	Q1'FY27	Margin
a. Radiopharma	136	16%	116	12%	121	12%
<i>Radiopharmaceuticals</i>	126	46%	106	33%	110	34%
<i>Radiopharmacies</i>	10	2%	11	2%	12	2%
b. Allergy Immunotherapy	63	35%	90	41%	66	31%
c. CDMO Sterile Injectables	62	17%	91	17%	45	9%
d. CRDMO	54	18%	64	20%	64	21%
<i>Drug Discovery Services</i>	32	20%	42	26%	45	26%
<i>CDMO – API</i>	22	15%	22	14%	19	14%
e. Generics	12	7%	32	15%	4	2%
f. Proprietary Novel Drugs	(6)		(7)		(8)	
<i>Unallocable Corporate (Expenses) / Income</i>	(18)		(22)		(25)	
Total EBITDA	302	15.8%	363	15.7%	268	11.9%

Other income for Q1'FY27 includes **grant income** of Rs. 5.6 Cr. for Line 3, which shall continue for more than 20 years and upto 30 years

Vision 2030

Revenue

Reach **2x** *from FY24 to FY30*

EBITDA Margin

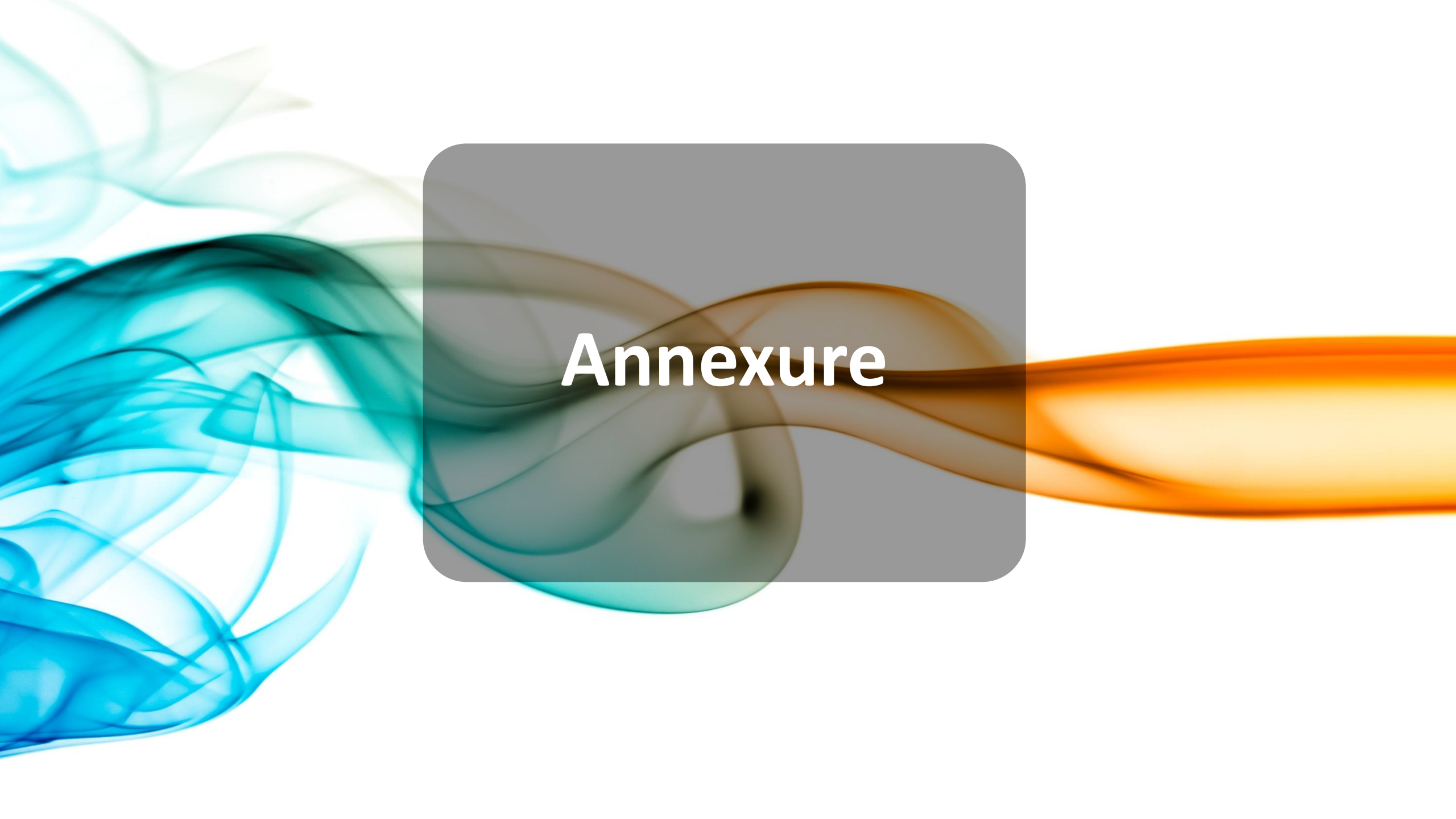
23% to 25% *by FY30*

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



Shantanu Jha
Group CHRO & Acting CEO,
Jubilant Biosys



Ashish Mukkirwar
CFO

Executive Leadership Team



Harsher Singh
CEO - Jubilant Radiopharma



Chris Preti
CEO - CDMO Sterile Injectables



Dr Jaidev Rajpal
CEO - Jubilant Generics



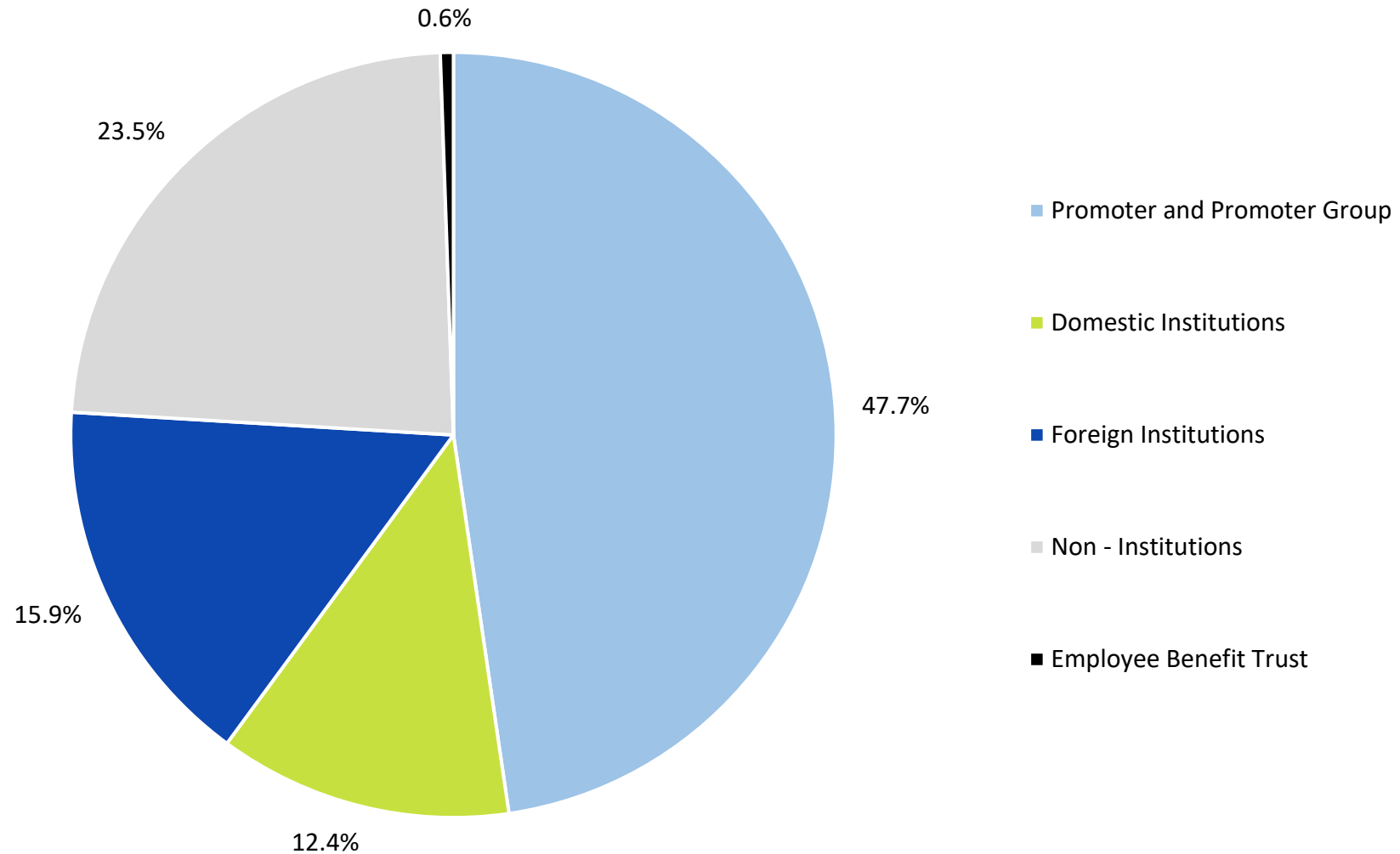
Kyle Ferguson
CEO - Allergy Immunotherapy



Daniel J. O'Connor
CEO – Jubilant Therapeutics

Shareholding Pattern

As on 30th June 2026



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 Inhibitor/Epigenetic Modulating Agent	CRISPR-Cas9 Endomorphic RNA Symptomatic Inhibitor (RNA based therapy targeting genetic disease)
PRMT5 Inhibitor	Medications that modify gene expression patterns
Brain Penetrant	Protein Arginine Methyltransferase 5 inhibitor (Blocks enzyme activity involved in adding methyl groups to arginine residues, affecting gene expression regulation)
PD-L1 Inhibitor	Cerebral blood flow enhancers or cognitive-enhancing drugs (supplements)
PAD4 Inhibitor	Programmed death Ligand-1 inhibitor (blocks the PD-L1 pathway, enhancing immune response against cancer cells)
LSD1/HDAC6 inhibitor	poly(ADP-ribose) polymerase 4 inhibitor (Disrupts DNA repair mechanisms in cancer cells, leading to their death)
NSCLC	Lysine specific demethylase 1/Histone deacetylase 6 inhibitor (Blocks enzymes involved in modifying histones, impacting gene expression regulation in cancer therapy)
SCLC	Non-small cell lung cancer
	Small cell lung cancer

A rack of test tubes containing liquids of various colors (blue, green, yellow, orange) with a semi-transparent dark grey bar across the middle.

Thanks!



Q1'FY27 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.

As we can demonstrate superior value proposition against competition, we are able to attract new channel partners. Our install base has grown by 26% in Q1'FY27 on an annualised basis. This improved scale is also helping to increase EBITDA margins in this product category.

We are also going to deploy 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'FY27?

Answer: The business continued to face unavailability of some SPECT products in Q1'FY27 as well. Post successfully conducting media fills, we have started commercial batch production at CMO Montreal in Q1'FY27. We expect first batches to be released in Q2'FY27 and production to normalize thereafter. We expect sales to normalize from H2'FY27 onwards.

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 are on-track for filing NDA in H2'FY27. We expect to launch MIBG 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 535 million. These products are generics or 505 B(2) versions of existing products and will be launched from FY28 to FY29. We got delayed in launching product in FY27, as we are now developing all our new products at a 3rd party CMO network, instead of CMO Montreal for risk mitigation. In addition to that, on the therapeutics side, we are working on MIBG.

Q5. Can you explain full year Q1'FY27 Radiopharmaceutical results?

Answer: Revenue grew 19% YoY to Rs. 322 Cr. on the back of sustainable & strong growth in Ruby-Fill ®. EBITDA for the period stood at Rs. 110 Cr. EBITDA margins decreased YoY due to unavailability of SPECT Radiopharmaceutical products at CMO Montreal. As supply resumes, Revenue and EBITDA will normalize from H2'FY27 onwards.

Q6. What is the impact of unavailability of SPECT products in revenue in Q1'FY27?

Answer: Q1'FY27 Revenue impact was approx. 7 Mn. Revenue is expected to return to normal levels from H2'FY27.

Q7. Can you update us on production situation for Radiopharmaceutical products. Can you talk about risk mitigation measures to ensure continuous supply for the business?

Answer: We are in good shape in terms of resuming the supply of SPECT products from CMO Montreal facility. We have successfully conducted Media fills in Q1'FY27 and have started the commercial batch production. We expect first batches to be released in Q2'FY27 and production to normalize thereafter.

On risk mitigation for future, we have ongoing technology transfer programs with both 3rd party CMO's and CMO Spokane for the SPECT products. We are tracking the schedule. It typically takes 18 to 24 months to complete the technology transfers. We shall gradually reduce the dependence on CMO Montreal.

Radiopharmacy

Q8. Can you talk about Industry demand? Where are we in the execution of new PET Manufacturing facilities?

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 and therapeutics. Notable ones include Life Molecular Imaging's F18 Neuraceq, Lantheus' F18 Pylarify and Novartis' Pluvicto.

We are investing USD 60+ million to expand our PET manufacturing 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 facilities shall be fully operational in FY28 and shall start to contribute significantly to the top line and bottom line.

Q9. Can you explain Q1'FY27 Radiopharmacy results?

Answer: Revenue grew 17% YoY to Rs. 700 Cr. on the back of increase in volume from certain PET products. EBITDA increased by 19% to Rs. 12 Cr. due to increase in revenue. Revenue from our current 3 PET manufacturing facilities continue to increase.

Allergy Immunotherapy

Q10. 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.

Q11. Can you explain Q1'FY27 Allergy immunotherapy results?

Answer: Revenues grew by 18% to Rs. 214 Cr. on the back of growth in revenue from the US market and outside US markets. EBITDA increased by 5% to Rs. 66 Cr. EBITDA margins are lower YoY due to lower production.

CDMO Sterile Injectable

Q12. 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 of higher-value products due to 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 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 to not only provide supply chain resiliency, but also to further mitigate any risk of new tariffs.

Q13. Can you talk about the launch of the third line at Spokane? What is the order book status and how do we see utilisations going forward? When can we expect launch of Line 4? What is the maximum revenue potential for Line 3 & 4 combined?

Answer: The capacity expansion program at our Spokane, Washington facility remains on track. Following the launch of our third Sterile Fill & Finish line (Line 3) in Q2'FY26, we are successfully ramping up revenues from technology transfer programs. Currently, 10+ products across multiple formats and vial sizes are undergoing technology transfer on Line 3. We are happy to share that we have onboarded one of

the world's largest oncology products on Line 3. Commercial batch production is expected to commence in late FY27, subject to FDA approval of these products.

Considering geopolitical uncertainties, large innovator pharmaceutical companies are increasingly seeking high-quality, US-based manufacturing capacities, specifically, significant capacities with isolator technology. As a result, we are seeing strong traction in Requests for Proposals (RFPs) for the new lines.

The next phase of capacity expansion—Line 4—is also complete. We expect this line to begin technology transfers by Q4'FY27 and then commercial production by end of FY28.

The revenue potential for Line 3 and Line 4 combined is estimated at USD 160 to 180 million. Also, we expect to see 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 at full utilisation.

Also, Please note that with Line 3 & 4, we are transforming ourselves from a small molecule CDMO to a specialized CDMO delivering complex biologics formulations for innovators. As we work with complex biologics, Customers demand specialized filling capabilities, tighter aseptic processing windows and stringent environment controls. These programs are difficult to onboard and longer to qualify, but once commercialized, they are more durable.

Q14: Can you talk about the severity of FDA observations at the recent audit at CMO Spokane?

Answer: FDA Audited CMO Spokane in Q1'FY27 and with 8 auditors over 8 days. There were no observations or comments related to sterility assurance concerns. We have responded to FDA and are addressing the observations comprehensively.

Q15. Can you give us an update on Montreal facility?

Answer: Construction work for the isolator-based new fill-and-finish line (Line 5) is progressing. The order for plant & Machinery has been placed and the steel construction shell is in place. The estimated total capex for the project is USD 114 million. Of this, approximately USD 35 million will be funded through concessional loans from the Canadian Government, with the remaining investment to be met through internal accruals. We expect installation to be completed by FY28, and technology transfer revenues to commence in FY29.

At the existing lines, the commercial batch production of radiopharmaceutical products has started, and the batches shall be released in Q2'FY27. The business will continue lean operations focussing initially on in-house Radiopharmaceutical products & then other customers. Post stabilisation of operations, we shall work to reduce EBITDA losses. Over the medium term, the new fill-and-finish line (Line 5) is expected to drive the growth & profitability of the business operations from FY29 onwards.

Q16: Is there any incremental impact post Warning letter at CMO Montreal? What kind of remedial actions are you taking? What is the financial impact of the same?

Answer: In Q1'FY27, the CMO Montreal received communication from USFDA conveying completion of review of EIR for inspection conducted from Oct20 to Nov3, 2025. The review resulted in issuance of Warning Letter.

We have sent a detailed response to FDA Warning Letter, in which we have supplied requested information and have given them update on status of independent third-party assessment on various areas. We are proactively engaged with FDA and will comprehensively address the concerns and observations.

The incremental remediation costs at the Montreal facility are primarily due to the need for additional external oversight, which is likely to continue into FY27.

Q17. Can you explain Q1'FY27 CDMO Sterile Injectables results?

Answer: Revenue grew by 34% to Rs. 496 Cr. due to incremental revenue from Line 3. EBITDA for the period stood at Rs. 45 Cr. EBITDA margins were lower YoY due to negligible third-party revenues & higher operating expenses including incremental remediation cost at Montreal facility.

Revenue at Spokane grew by 43% to Rs. 494 Cr. and EBITDA grew by 49% to Rs. 117 Cr. EBITDA margins also expanded by 100 basis points to 24%.

CRDMO – Drug Discovery

Q18. Can you talk about demand scenario in Drug Discovery services? How do you see revenue 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 Biosecure ACT, which was enacted into law in Dec'25. We are increasing our partnership with large Pharma companies, leveraging our infrastructure, capacity and capabilities expanded during last two years.

We are well prepared to further scale up Infrastructure and scientific talent to take advantage of the upcoming in CRO demand. We have talked about increasing our FTE capacity to 4,000 FTEs in phased manner to cater to increasing demand. We expect a healthy revenue growth to continue along with steady margins.

In the short term, we expect competitive intensity to increase in the large-pharma customer segment, while demand conditions in the biotech segment are expected to improve.

Q19. Can you explain Q1'FY27 CRDMO Drug Discovery results?

Answer: Revenue grew by 8% to Rs. 174 Cr. EBITDA for the period grew by 43% to Rs. 45 Cr. due to operating leverage and forex gain.

CRDMO – API

Q20. Can you give us update on custom manufacturing business?

Answer: In Q1'FY27, Revenue mix from custom manufacturing has gone up on YoY basis.

Q21. Can you explain Q1'FY27 CRDMO API results?

Answer: Revenue for Q1'FY27 stood at Rs. 135 Cr. EBITDA for the period stood at Rs. 19 Cr. Revenue and EBITDA margins decreased YoY due to the industry wide pricing pressure and RM cost increase due to middle east conflict. We are consciously moving the revenue mix towards profitable products. Going forward, we expect the custom manufacturing revenue mix to drive the utilisation.

Generics

Q22. Can you tell us your plans for new product launches?

Answer: Since April'24, we secured approval of (12) ANDA's from our pipeline. We have launched 4 new products in our US last year. We have launched 2 new products in Q1'FY27. In the current year, we expect to launch 6 to 8 products in the US and Non-US market. Therefore, we have an improving growth and profitability outlook.

Q22. Can you explain Q1'FY27 generics results?

Answer: Revenue grew by 4% to Rs. 173 Cr on the back of launch of 2 new products. EBITDA for the period stood at Rs. 4 Cr. EBITDA margins decreased YoY due to change

in product mix. Trailing 12 months margin remains at ~10%. Expect margins to increase in subsequent quarters.

Prop Novel Drugs

Q24. 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 I clinical data preliminary demonstrated a manageable safety profile 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 findings, we have initiated a Phase I/ II clinical trial to treat ET and MPN patients with Thrombocytosis (high platelet counts). The study is ongoing and preliminary data is showing rapid and durable dose dependent platelet normalization in patients with Essential Thrombocythemia (ET) and MPNs in Australia.

The initial phase I trial in the US also showed anti-tumour response in two lung cancer patients. One non-small cell lung cancer (NSCLC) patient with STK11 mutation, having progressed on prior doublet immuno-oncology (IO) therapy, showed anti-tumour activity. Generally, the survival rate is very low in such cases, however, the patient has responded well to JBI-802 monotherapy. Therefore, an investigator led clinical trial in NSCLC has been initiated and is ongoing 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 1, first in human study, for this molecule, in patients with specific cancer sub-sets at major oncology centres in India.

Consolidated Financials

Q25. Can you talk about overall financial performance in Q1'FY27?

Answer: In Q1'FY27, Revenue grew by 17% on a YoY basis to Rs. 2,229 Cr. on the back of growth across all Business units. EBITDA for the quarter stands at Rs. 268 Cr. EBITDA margins decreased YoY primarily due to unavailability of high margin SPECT Radiopharmaceutical products and negligible third- party revenues & higher operating expenses including incremental remediation cost at CMO Montreal. Other income for Q1'FY27 includes grant income of Rs. 5.6 Cr. for Line 3, which shall continue for more

than 20 years and upto 30 years. Reported PAT for the period stands at Rs. 56 Cr. PAT margins decreased due to lower operating profitability and increase in depreciation.

As we are consciously investing in businesses to secure future growth, Net Debt / EBITDA increased from 1.5x in Mar'26, to 1.8x in Jun'26.

Q26. What is the outlook for FY27?

Answer: We expect growth momentum to continue from Q1'FY27 to rest of the year. In terms of EBITDA margins, it shall be story of two halves. As production at CMO Montreal stabilises, we expect EBITDA margins to strengthen in H2'HY27.

In terms of capex, we expect to have a similar outlay as in FY26.

Q27. Can you talk about the impact of US tariffs on the business?

Answer: Jubilant Pharmova Limited derives approximately 81% (Q1'FY27) of its revenue from the US market. It is therefore imperative to note the implications of the multiple new tariffs announced by the US government on the company's various business segments.

The origin of the goods and services sold in the US by the Company (Q1'FY27) is approximately 78% from the US itself, 14% from Canada and 8% from India.

The goods and services originated and sold in the US itself are mainly from Radiopharmacy business, Allergy Immunotherapy business and CDMO Sterile Injectable business. Among these three businesses, the company continues to have strong positive impact on its CDMO Sterile Injectable business. The business primarily manufactures innovator products and has large innovator companies as its customers. Due to the new tariffs, the large innovator companies are now looking to create an alternate manufacturing site in the US, for their US requirements. This has led to an excellent traction in RFP's and order booking for the Company's new Line 3 in Spokane, Washington.

The goods and services originated in Canada and sold in the US are 14% (Q1'FY27) of the Company's US revenue. The goods exported from Canada include Radiopharmaceutical products, which are exempted from tariffs under US, Canada and Mexico trade agreement. Therefore, this business will have no material negative impact.

The goods and services originated in India and sold in the US are 8% (Q1'FY27) of the Company's US revenue. The goods exported from India include Generic finished

formulations and Generic Active Pharmaceutical Ingredients (APIs) products. As a risk mitigation strategy, in the generics finished formulation business, the company has also developed CMO network through partners with facilities in the US as well.

In summary, the company expects overall positive impact of these new US tariffs, especially on its CDMO Sterile Injectable business with no material negative impact in rest of its business segments.

.....*End*