JUBILANT PHARMOVA Investor Presentation Jun'24





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 - Snapshot



Jubilant Bhartia Group founded by Shyam S. Bhartia and Hari S. Bhartia, leading industrialists from India





Strong presence in diverse sectors like Pharmaceuticals, Life Science Ingredients, Contract Research & Development Services and Therapeutics, Performance Polymers, Food Service (QSR), Food, Auto, Consulting in Aerospace and Oilfield Services



Global presence through investments in India, USA, Canada, Europe, Singapore, Australia, Africa, China, Sri Lanka and Bangladesh



Employs around 46,000 people across the globe with ~2,200 in North America

Company Snapshot



A global pharmaceutical company with strong team of approx. 5,500 multicultural people & Revenue at Rs. 6,703 Cr. (FY24)



Revenue Split (BU wise)





Global Manufacturing & Research Footprint



World class manufacturing facilities, 2 state of the art research centers & 46 radiopharmacies

Kirkland, Montreal, Canada CDMO – Sterile Injectables



Kirkland, Montreal, Canada Radiopharmaceuticals





Spokane, Washington, USA CDMO – Sterile Injectibles Spokane, Washington, USA Allergy Immunotherapy





INDIA

NORTH

AMERICA



Roorkee, Uttrakhand, India Generics





Nanjangud, Karnataka, India API

G. Noida, Uttar Pradesh Drug discovery, CDMO



Chemistry Innovation Research Center Greater Noida



Bengaluru, Karnataka Drug discovery

Jubilant Pharmova in FY24



Achieved strategic milestones; Improved overall financial performance to drive shareholder value



FY24: Revenue (+ 7% YoY), EBITDA (+20% YoY), Net Debt / EBITDA (Reduced from 2.93x to 2.48x)



Growing role in treatment of life threatening diseases

- Radiopharmaceutical is a combination of radioactive isotope and pharmaceutical drug
- Radiopharmaceuticals are used to diagnose and to treat life threatening diseases e.g. Cancer, Cardiac disorders, Neurological disorders
- There are **3 type of** ٠ procedures that use radiopharmaceuticals
 - **SPECT Imaging** •
 - **PET Imaging**
 - **Therapeutics**

	Single-photon Emission Computed Tomography (SPECT Imaging)	Positron Emission Tomography (PET Imaging)	Radiopharmaceutical Therapeutics (Tx)
Description	 Uses "low-energy" radio isotopes that emit gamma rays, detected by SPECT cameras 	 Uses "high energy" radio isotopes that emit positrons, detected by a PET scanner 	 Radiation is systemically or locally delivered using pharmaceuticals that either bind preferentially to targeted cells or accumulate physiologically
Key Facts	 Longer half-lives Images blood flow Specialized but legacy products, > 90% generics 	 Shorter half-lives Images blood flow and metabolic processes Superior image quality Mostly innovative, few generics 	 Specialized / new generation isotopes Targeted therapies with higher efficacies Minimal off target toxicity vs. conventional treatments
Market trends	 Large and Stable market Robust supply chain management 	 High growth market More expensive vis-à-vis SPECT 	 High no. of clinical trials in the space Accelerating M&A activity in therapeutics space with multiple > USD 1 Bn. deals in 2023
Key Products & Isotopes	 MAA, DTPA, Exametazime, Sulfur Colloid, Mertiatide Isotopes - Tc99 	 Ruby-Fill ®, Pylarify, Illuccix, Neuraceq, FDG Isotopes - Rb82, F18, Cu64 	 Products - HICON® Sodium Iodine I 131, Pluvicto, Lutathera Isotopes - Lu177, Ac225, Pb202
Mode			

of Operation









US radiopharmaceutical market is expected to reach approx. USD 20 Bn. by 2030, growing at a CAGR of 20 %+

US Radiopharmaceutical Market (USD Bn.)



Growth Drivers and Key Trends

- Growth driven by superior imaging and therapeutics profiles, new emerging isotopes with low off target toxicity and increasing use cases for un-met needs
- PET imaging market growth is fueled by novel products, e.g., PSMA sales has exceeded USD 1 Bn. in <2 years of launch. PET market growth is driven by
 - Strong fundamentals such as better imaging, significantly lower false negatives and faster examination time
 - Applications extending beyond oncology, such as Cardiology scans, Alzheimer's
- Advanced Radiopharmaceutical Therapy market is witnessing launch of differentiated, high value and high efficacy products e.g. Pluvicto used for Prostate Cancer exceeding USD 1 Bn. sales.
 - Favorable pharmacological profile with lower toxicity and higher efficacy, especially in areas with un-met needs
 - New / emerging isotope profiles with targeted effects and lower off target impacts, such as Lu177 and Ac225
 - Application in therapeutic areas beyond oncology such as Neurological conditions, e.g. Alzheimer's

Consolidated market with high entry barriers



Stringent manufacturing & regulatory environment

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

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

Innovative new product development

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

Forward integration with radiopharmacies

Forward integration with radiopharmacies helps to gain market share



We are one of largest manufacturer in the addressable market in the US with a wide radiopharmaceutical portfolio

Organ	Туре	Product	Key Indication		
Luna	SPECT	Tc99m-DTPA	Pulmonary Embolism		
Lung	SPECT	Tc99m-MAA	Pulmonary Perfusion		
Thyroid	SPECT	I-131	Localizing metastases associated with thyroid malignancies		
	Тх	I-131 HICON®	Hyperthyroidism, Selected cases of Carcinoma of Thyroid		
	PET	Ruby - Fill ®	Coronary Artery disease		
Cardiac	SPECT	Tc99m-Gluceptate	Cardiac blood pool Imaging		
	SPECT	Tc99m-Sestamibi	Coronary Artery disease		
Breast	SPECT	Sulfur Colloid	Localization of metastatic lymph nodes, imaging of liver, spleen		
Gastrointestinal	SPECT	Tc99m-Exametazime	Intraabdominal Infection		
Renal	SPECT	Tc99m-Mertiatide	Renal failure, Urinary tract obstruction		
Muscoskeletal	SPECT	Tc99m-MDP	Delineate areas of altered osteogenesis		

Key Differentiators

- Diversified product portfolio spread across SPECT & PET diagnostics and growing therapeutics
- **High profitability** owing to efficient cost structure, in-house APIs and robust supply chain management
- Partner of choice for leading customers owing to innovative products with superior profile vs. competitors and best in class customer service, e.g. Proprietary Ruby-Fill ® technology for Cardiac Imaging
- On-shore manufacturing facility in Montreal with high quality track record and ability to manage complex processes
- Strong R&D capabilities, continuously feeding the product pipeline to enable frequent market launches

Market leadership in select products - MAA, DTPA and I-131

Draximage ® MAA



MAA is used in the perfusion phase of a ventilation/perfusion (V/Q) scan to diagnose pulmonary embolism. JDI is market leader 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 the sole supplier 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



HICON® Sodium Iodine I 131 Solution USP





Innovation Leadership in Ruby - Fill [®] which is gaining market share consistently

Growing Cardiac PET Market in the US



Source : Company Estimates

Growth Drivers and Key Trends

- Superior product profile vs. SPECT scans
- Improved reimbursement landscape and diagnostic infrastructure
- Lower half lives vs. SPECT products leading to lower hospital burden

Ruby-Fill ® Rubidium 82 generator and Elusion System

- The RUBY-FILL® Rubidium 82 Generator contains accelerator produced Strontium-82, which decays to Rubidium-82 (Rb-82). It is used for Cardiac PET scan, a non-invasive imaging procedure of the myocardium, to evaluate regional myocardial perfusion in adults with suspected or existing coronary artery disease.
- Ruby-fill is installed in top 80% US Cardiac networks and is positioned to further increase market share
- Lower cost vs. competition driven by higher shelf life of generators, hence driving more scans per generator
- Better Image quality due to patented feature of saline push dosing, significantly increasing Rb-82 activity delivered to the heart
- Consistent image quality due to proprietary constant activity mode, plus patient weight based dosing

Gaining market share in the US cardiac PET market; Record Ruby-Fill® installations in the US in FY24

Ruby-Fill[®] and Robust product pipeline to fuel future business growth





Ruby-Fill[®] Growth potential

- Gain market share in the growing US cardiac PET market
- Scale ex-US markets such as Europe, Canada, etc.



PET & SPECT Product Pipeline

- Target to launch new products in PET Imaging with an addressable market at ~ USD 500 Mn.
- Pipeline in SPECT Imaging with an addressable market at ~ USD 50 Mn.



Development of therapeutic product - MIBG

 Completed patient dosing for Phase II clinical trials for MIBG. Expect launch for relapse / refractory Neuroblastoma (~400 patients per annum) in CY 2026.

Driving revenue growth



Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	215	241	256	19%	872	952	9%
EBITDA	100	126	126	26%	465	477	3%
EBITDA Margin (%)	47%	52%	49%	250 bps	53%	50%	(320) bps

- FY24 revenue grew YoY on the back of **new products sales** in **Mertiatide**, **Sulfur colloid** and growth in **Ruby-Fill**®
- FY24 EBITDA increased YoY on the back of increase in revenue



US Radiopharmacy market is expected to grow on the back of novel PET & Therapeutic products

SPECT Radiopharmacy



PET Radiopharmacy





Growth Drivers and Key Trends

- Radiopharmacy dispenses and distributes radiopharmaceutical products
- Consolidated market in US with top 3 radiopharmacy networks dispensing and distributing 70%+ products
- Increasing demand of novel PET diagnostics product, e.g., Cyclotron based pharmacies for F-18 PSMA, Alzheimer's products. Additionally, SPECT pharmacies can handle generator based PET products, e.g., Ga-68 PSMA
- Therapeutics dispensing share of pharmacy networks expected to grow, driven by Stringent USP 825 regulations. Most clinics and hospitals don't want to invest in the clean room infrastructure for dispensing. Additionally, big pharma companies have limited capabilities in the distribution and handling wastes of radioactive materials
- Emerging radioisotopes landscape such as Rb-Sr, Ga-68, Cu-64, Lu-177, Ac-225 are leading to development of new PET Imaging and Theranostic products which will further fuel radiopharmacy share of dispensing and distributing these products. 16

1. USP develops uniform minimum standards for the preparation, compounding, dispensing, and repackaging of radiopharmaceuticals





Consolidated Market

	# of radio pharmacies in the US	SPECT pharmacies	PET pharmacies	# of hospitals served in the US
Cardinal Health	160+	✓	✓	~ 4,100
	46	✓	✓	~ 1,800
SIEMENS Healthineers PETNET Solutions	41		✓	~ 700
🔯 RLS	31	✓		~ 900
Pharmalogic Take The Lead	42	✓	✓	~ 200
	14		✓	~ 200

Barriers to Entry



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



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Intricate Supply Chain

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

Complex Care Coordination

Requires awareness, education, and collaboration across multiple hospital departments

Skilled Manpower Requirement



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



2nd largest radiopharmacy network in the US





Ride on volume & new product led industry growth, evaluate opportunities for network expansion





Strategic investment in PET radio pharmacies yielded solid returns & validated our investment thesis

Value Creation by Investment in PET radiopharmacy Business



Validation of our Investment thesis in PET radiopharmacy Business

- JPL, Company's wholly owned subsidiary invested USD 25 Mn. in Nov'2020 in Sofie Biosciences Inc. ('Sofie'). JPL holds 25.8% stake
- Sofie entered in a definitive merger agreement with Trilantic Capital Partners, North America, a US private equity firm.
- Consequently JPL planed to sell its entire 25.8% equity stake in Sofie for aggregate proceeds of about USD 142.9 Mn. (including preferred returns). Of this, USD 115.9 Mn has been received by JPL by Jun'24. An additional USD 1.2 million is held in escrow account for any post-closing adjustments. Balance sum of upto USD 25.8 Mn. is contingent upon achievement of certain future milestones.
- Plans to use funds to reduce debt, capex and other corporate purposes

USD 115.9 million proceeds received by JPL by Jun'24



Volume to drive revenue growth & operational efficiency to drive margin expansion

Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Ү-о-Ү
Revenue	475	511	561	18%	1,681	2,050	22%
EBITDA	(4)	10	38	995%	(87)	56	164%
EBITDA Margin (%)	(1%)	2%	7%	760 bps	(5%)	3%	790 bps

- FY24 revenue grew YoY on the back of increase in volume from new products
- FY24 EBITDA increased YoY on the back of increase in volume & improvement in operational efficiency

> 50 Mn.

Americans suffer

from some type

of an allergy

annually

Global market poised to reach USD 3 Bn. by 2028, growing at a CAGR of ~ 7%

Most Common Allergies in US (2023)



Global Allergy Immunotherapy Market (USD Bn.)



- Allergy immunotherapy (AIT) refers to the treatment for allergic reactions against a variety of allergens including Pollen, Mold, PET dander, Food & Insect (treated by venom immunotherapy) etc. In this treatment, repeated shots of allergic antigens are provided to develop immunity and eventually cure allergy over a period of time.
- There are two kinds of delivery mechanisms Sub Lingual and Sub Cutaneous

Growth Drivers

>50

Deaths in US in a

year due to

Anaphylaxis

- Increasing allergy cases
- Awareness of allergy treatment
- Advancement in treatment options



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Jubilant is #2 player in the US Sub-Cutaneous Allergy Immunotherapy market with strong entry barriers

Strong Entry Barriers

- Highly concentrated US market with well established players
- Raw material comprising natural extracts / organisms involve a complex supply chain from sourcing to processing.
- Grandfathered approvals with any new product needing a Biologic License Approval which is more complex than small molecule drug approval.
- In order to succeed, New Entrant has to offer a complete portfolio of products, which shall entail significant investment, development and approval lead times.

Key Differentiators

- # 2 player in the US SCIT allergy market & Sole Supplier of Venom immunotherapy in the US since 2018.
- Product portfolio includes 6 different Insect Venom products, 200+ allergenic extracts and skin testing devices, with best in class customer service and high supply reliability.
- **HollisterStier'** brand loyalty going back 100 years
- Onshore USFDA approved Manufacturing. Dedicated Sales force in the US, serves over 2,000 customers including Allergists, ENT Physicians

Balanced Product Portfolio



Venom Extracts



- Venom extracts includes products for Honey Bee, White-Faced Hornet, Yellow Hornet, Wasp, Yellow Jacket and Mixed Vespid allergies
- Sole supplier in US

Allergenic Extracts



- Allergenic extracts (over 200 products) includes products for Dog, Cat, Mite, Tree Pollen
- Combination of specialized (e.g., Dog) and standardized extracts (e.g., Cat); 2nd largest in the US

Skin Testing Devices



- Multiple skin test system includes ComforTen, Quintest and Quintip
- Differentiated product vs. competition – stainless steel lancets vs. plastic tips ensuring minimal trauma

Moving ahead on three pronged growth strategy





Enlarge US Venom Segment

- Create customer awareness on the Bee sting allergy through targeted marketing campaigns and enlarge the US Venom segment
- Leverage Brand equity in the community



Gain market share in US Allergenic extracts

- Use Venom products to gain customer wallet share in Allergenic extracts
- Launch differentiated products e.g. Ultra Filtered Dog product



Penetrate outside US market

- Penetrate the Europe market on the back of **strategic partnerships**
- Expand the distribution channel in APAC, MEA & LATAM

Sustained growth momentum & margin expansion



Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	170	161	188	11%	603	679	13%
EBITDA	55	62	75	36%	206	273	33%
EBITDA Margin (%)	33%	38%	40%	750 bps	34%	40%	620 bps

• FY24 revenue grew YoY on the back of volume & price increase

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FY24 EBITDA margin increased YoY due to increase in revenue and improvement in operational efficiencies

3 CDMO - Sterile Injectables Demand expected to outpace the supply by FY'26



Global CDMO-SI Market Size (in USD Bn.)



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.

Growth Drivers & Key Trends

- Increase in demand: Increasing number of drugs/injectable in development pipeline driven by biologics (65%+ of current pipeline) and LOEs
- Increase in outsourcing: Outsourcing expected to increase, driven by limited internal capacity and capabilities, cost reduction initiatives and big pharma focus on internalizing specialized capabilities, e.g., Gene Therapy, Peptides
- Significant shortages: Since 2015, 50-60% of new drug shortages in the US have been injectables, signaling need for significant on-shoring
- Demand Supply Gap expected to widen further with increasing consolidation, e.g., Novo Holding acquired Catalent for enterprise value of USD 16.5 Bn. This transaction may further reduce the overall capacity available for outsourcing, given Novo is expected to use capacity for manufacturing their anti-obesity drugs.

Structurally attractive market with key differentiators driving our growth

Strong 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 (e.g., Multi Dose Preservative Free ophthalmic drops, etc.) have emerged, driven by requirements of differentiated technologies, 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

Key Differentiators

- Deep and long-term relationships with our customers
 Top 10 Customers have been with us 5+ years.
- On Shore Manufacturing facilities in Spokane, US and Montreal, Canada
- Co-invested capacity with US govt., advanced isolator technologies are part of our expansion, meeting both regulatory & customer requirements
- Steady quality track record in past audits including inspections from US FDA, ENVISA Brazil and others
- Focused core competency in Sterile Fill & Finish and Ophthalmic (ointments, liquids & creams) sterile products
- Customer satisfaction is strong with 90%+ repeat Customer business rate





Collaborative partner with unique capabilities & strong customer relationships



Full Suite of Services with On-shore manufacturing

Strong Quality track record

- Can handle Vial size from 2ml to 100 ml with batch size up to 2,000 ltr.
- **Full suite of services** including sterile fill and finish (Liquid & Freeze dried), Ophthalmic (Liquids, Ointments and creams) and Biologics
- Strategically located on-shore manufacturing footprint in North America

- Steady quality track record in past audits at each location including inspections from FDA, ENVISA and others
- Stability in core portfolio at Spokane with **multiple products having patent protection** and limited competition

Strong Customer Relationships

- Serve leading pharmaceutical companies globally
- Long standing relationship with customers with some longer than10 years and 90%+ repeat business rate
- Customer-focused approach with strong Tech Transfer & Project Management collaboration from the development phase



Doubling of capacity with state of the art technology at Spokane on track





- Doubling Spokane capacity of sterile fill and finish (both liquid and lyophilization) within stipulated timelines and costs. Total investment at USD 285 Mn. partly funded through cooperative agreement with US Govt. for USD 149.6 Mn.
- Line 3 commercialization expected in FY'26, followed by Line 4 in FY'28

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Reverted to normalised operations in Q4'FY24

02/5/24

Particulars (RS. Cr.)	Q4 F125	Q3 F124	Q4 F124	1-0-1		F125	F124	1-0-1
Revenue	321	303	259	(19%)		1,155	1,117	(3%)
EBITDA	86	37	58	(32%)		345	192	(44%)
EBITDA Margin (%)	27%	12%	22%	(430) bps		30%	17%	(1,270) bps
Adjusted Revenue		1,063	1,117	5%				
Adjusted EBITDA		258	192	(26%)				
Adjusted EBITDA Mar	gin	24%	17%	(710) bps				

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shutdown and impact of **COVID related business**

FY24 EBITDA decreased YoY due extended

to extended shutdown in Q3'FY24

Adjusted EBITDA decreased in FY24 due to proactive remediation and planned extended shutdown in Q3'FY24



4 CRDMO: Drug Discovery Services, CDMO & API



Both Drug Discovery Services and API/Formulation Development markets are expected to grow at ~5-6% CAGR



API/Formulation Development Market Size (USD Bn.)



Growth Drivers for Drug Discovery Market

- Large Pharma companies to de-risk their supply chain by adding "friend sourcing" locations. Biosecure Act is proposing to prohibit US Govt. and US life sciences companies, (who are receiving federal grant money) to work with biotechnology service providers that are connected to foreign adversaries
- Rise in **specialized discovery technologies** such as ADCs and oligonucleotides
- Lower Biotech funding stalled growth, however early signs of recovery with further recovery expected by late FY'25

Growth Drivers for API / Formulation development Market

- Focus on integrated service offering ranging from discovery to development
- Rapid momentum in specialized CDMO services to support ever increasing clinical trials, e.g., High potency APIs with stringent exposure control requirements
- Rising share of biologics along with increasing investments in biologics for new niche modalities

Source : Company Estimates

CDMO API market



CDMO API Market is estimated to grow at a CAGR of ~ 6%+ from 2020 to 2026



Growth Drivers for API Market

- Although API market is dominated by the small molecules, higher growth is experienced by HPAPIs and large molecule segments
- Cost competitiveness is the key including backward integration into major KSMs to mitigate pricing pressures on the finished good formulation companies
- Rising interest of companies in manufacturing **custom generics** for innovators, ensuring higher margins
- Move towards friend sourcing becoming increasingly apparent, reducing concentration risk of generics API manufacturing

CRDMO : Drug Discovery Services & API

We provide end to end CRDMO services for drug substance in small molecules





Drug Discovery Services

Three Pronged growth strategy





On boarded two large pharma clients in FY24; well prepared to scale up infrastructure (labs, scientific talent etc.) to take advantage of increase in CRO demand



Maximize market penetration & Transform operations by increasing cost effectiveness & asset utilisation



State of the art GMP manufacturing facility spanning over 41 acres with 7 multi stream manufacturing blocks

Facility inspected by FDA, PMDA Japan, KFDA Korea, ANVISA Brazil, TGA Australia

Dominant position in select therapies

- Comprehensive portfolio comprising of APIs from various therapeutic area -Central Nervous System, Cardiovascular System, Anti-infective and Antidiabetic
- Among the largest producers for API's such as, Oxcarbamazepine, Carbamazepine, Pinaverium, Resperidone, Donepezil, Lamotrigine, Meclizine, Azithromycin & Valsartan
- Reach to 50 countries, Servicing 160+ customers

Strategy going forward

- Maximize penetration: Fortifying sales in USA, Japan, LATAM, MENA regions
- Transform operations: Increasing overall cost effectiveness & asset utilisation
 - Increase backward integration: De-risk by increasing backward integration & follow China plus one strategy for sourcing
 - Increase capacity utilization: Leverage GMP manufacturing capabilities for Innovative APIs (CDMO)



CRDMO API: Pricing pressure continues; Taking initiatives to reduce operating costs & increase capacity utilization

Drug Discovery Services

Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	131	114	117	(11%)	522	449	(14%)
EBITDA	35	30	29	(18%)	164	106	(35%)
EBITDA Margin (%)	26%	27%	24%	(200) bps	31%	24%	(780) bps

API

Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	163	138	165	1%	663	645	(3%)
EBITDA	12	11	24	101%	35	63	80%
EBITDA Margin (%)	7%	8%	14%	720 bps	5%	10%	450 bps

- FY24 revenue decreased YoY. Industry headwinds in Biotech Industry is on account of lower funding for early stage drug discovery projects. **On boarded two new large Pharma clients.**
- FY24 EBITDA decreased YoY on account of reduced revenue.

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- **FY24 revenue marginally decreased** due to pricing pressure in certain products.
- Q4'FY24 and FY24 EBITDA increased YoY significantly due to cost optimization

CRDMO Segment

Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	294	252	282	(4%)	1,185	1,093	(8%)
EBITDA	47	41	52	13%	199	169	(15%)
EBITDA Margin (%)	16%	16%	19%	280 bps	17%	15%	(130) bps









■ Developed Markets ■ Pharmerging Markets ■ RoW Markets

Overall Market

 Overall market is growing on the back of increase in Chronic disease prevalence, loss of exclusivity for innovator products negated by pricing pressure in select markets

Developed Market

- US market is expected to grow ~2% with early signs of decrease in price reductions. Legacy generics expecting price reduction of approx. 7% in FY24 vs 12% in FY23
- Non-US market is expected to grow by ~5 to 7% with margins & regulatory approval timelines varying by market. Key differentiators are cost competitiveness and supply reliability.

India Market

 India market is expected to grow in excess of 10%. Key differentiators are brand building and In-clinic effectiveness of sales team

Generics



Target to reach EBITDA breakeven in short term & then grow profitability in the medium term

Key Products & Facilities

- Therapeutic areas Cardiovascular System, Central Nervous System, Gastrointestinal and Multi Specialty
- Global presence with serving more than 50 countries including US, Europe, Canada, Japan, Australia and RoW
- Building branded generics business in India in the field of Cardiovascular diabetes & Multi Specialty
- **Derisking product supplies** through building a robust CMO network
- USFDA classifies Roorkee Facility as "Voluntary Action Indicated (VAI)" in April'24.



Generics Growth Strategy for key markets





Grow the profitable Non-US International market

- Focus on scaling 2 key markets to triple digit revenue in INR Cr. (B2B2C)
- Offer a portfolio of products to 50+ markets (B2B)
- Launch new products through In-Licensing





- Build and Scale branded
 generics business in India
- Develop 3 to 4 profitable therapeutic area divisions. Demonstrated successful blueprint by achieving profitability in CVD division in Q4FY24



Achieve profitability in the US Market

- Focus on profitable sustainable portfolio
- **Outsource** manufacturing to CMO's. Launch **new products**
- Relaunch products & grow exports through Roorkee Facility



Generics

Actions to propel US Generics business towards profitability



Generics







Continuous Quality Improvement

Implemented a large scale **quality improvement** program in Roorkee facility.

Continue the upgrade the quality framework



De-risk Product supplies by outsourcing

De-risking product supplies through building a **robust CMO network** & outsource the manufacturing

Wide network of CMO's being built across US, Europe, India and other countries



Continue Cost Optimisation

Implemented cost optimization initiatives of **Rs. 150 Cr.**

Continue to implement cost saving opportunities

Generics *Moving in the right direction*



Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	199	199	201	1%	762	775	2%
EBITDA	(39)	(31)	(39)	1%	(230)	(141)	39%
EBITDA Margin (%)	(20%)	(15%)	(19%)	30 bps	(30%)	(18%)	1,210 bps
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Adjusted Revenue	734	799	9%
Adjusted EBITDA	(241)	(104)	57%
Adjusted EBITDA Margin	(33%)	(13%)	1,980 bps

- Target to reach EBITDA breakeven in short term
 - Grow profitable Non-US international business
 - Achieve profitability in the US business
- Grow profitability in the medium term
 - Relaunch products & increase export through Roorkee facility in a gradual and meaningful manner

6 Proprietary Novel Drugs Clinical stage precision therapeutics



Advancing potent molecules to address unmet medical needs in Oncology & Auto immune diseases

Program	Mechanism	Indications	Lead Optimization	Pre - Clinical (IND)	Phase I /II	Milestones
JBI-802	coREST Inhibitor/ Epigenetic Modulating Agent	ET(Essential thrombocythemia)/MPN (Myeloproliferative			0	Phase I data suggests therapeutic potential. Early Phase II data in
		small cell lung cancer)				ET / MPN in H2-2024
JBI-778	PRMT5 Inhibitor Brain Penetrant	EGFR (Epidermal Growth Factor receptor) refractory NSCLC, ACC (Adenoid cystic carcinoma), High Grade			0	Phase I / II initiation in H1 2024
		Glioma				
JBI-2174	PD-L1 Inhibitor Brain Penetrant	Brain tumor and metastases		0		IND enabling
JBI-1044	PAD4 Inhibitor	Oncology and auto-immune disease		0		IND enabling
Other	Various	Various	—0			Undisclosed Research Programs

Two Clinical stage drugs under development with significant value inflection potential on clinical outcome

Key Indications for JBI - 802



Disease Indications	Rationale	JBI - 802 Response
Non-Small cell lung cancer (NSCLC)	 STK11 mutation is observed in10-15% NSCLC (85 % of lung cancer is NSCLC). Patients with STK11 mutations have a lower survival rate and are resistant to immune checkpoint therapy (like Keytruda, Atezolizumab, etc.) 	One patient with NSCLC having STK11 mutations showed significant response on JBI-802, while not responding to previously administered doublet Immune checkpoint therapy. Preclinical animal model study have shown synergistic effects of JBI-802 with immune checkpoint inhibitors
Essential Thrombocythemia (ET)	 ET is a rare blood cancer that causes the bone marrow to produce too many platelets which can lead to an increased risk of developing blood clots resulting in stroke and heart attack Limited options for patients who are refractory to the first line of therapy 	JBI-802 has shown to reduce platelet in human clinical trial which is mediated by LSD1 inhibitor. JBI-802 has better safety profile compared to the competitor (no Dysgeusia and anemia)
Post MPN-AML (Myeloproliferative neoplasms- Acute myeloid leukemia)	 MPNs are a group of blood cancers that cause increased production of blood cells, mainly affecting red blood cells, platelets, or white blood cells. Progression from MPN to AML (Acute Myeloid Leukemia) is a serious complication, occurring in about 5-10% of MPN patients. No effective therapy available (Survival in adults is only 5 months) 	JBI-802 shows superior efficacy in preclinical in-vivo efficacy studies compared to LSD1 only and HDAC6 only inhibitors

Phase Two & Investigator led clinical trials to start for JBI-802

JBI-802 alone or in combination with an α-PD-1 monoclonal antibody in CT-26 Syngeneic Model

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JBI-802 in combination with α-PD-1 achieves tumor regression Complete tumor regression achieved in 3/8 animals

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Product

JBI-802 Small molecule - CoREST Inhibitor

Indications

- Essential Thrombocythemia / Myeloproliferative Neoplasms
- Lung cancer (NSCLC), Acute myeloid leukemia (AML)

Scientific Rationale

- CoREST inhibition by dual targeting LSD1 and HDAC6
- Superior preclinical efficacy vs other LSD1 or HDAC6 single agent inhibitors
- Dual targeting mechanism Synergistic efficacy
- Anemia not seen in both preclinical and clinal studies, Dysgeusia not seen in clinic v/s other drugs

JBI-802, LDS1i or HDAC6i in HEL 92.1.7 Xenografts

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JBI-802 observed to have superior anti-tumor activity vs inhibition of LSD1 or HDAC6

Development pathway

- Phase 1 clinical data establishes safe dose and showed anti-tumor response in 2 lung cancer patients at the low dose of 10mg without platelet reductions
- One patient with NSCLC having STK11 mutations showed significant response on JBI-802, while not responding to previously administered doublet IO therapy
- Also dose dependent platelet effect seen in clinic, establishing application in Essential Thrombocythemia (ET) and other Myeloproliferative Neoplasms (MPN)
- Phase II clinical trial to treat ET and MPN patients with thrombocytosis is being initiated in H1 2024
- *Investigator led clinical trials* in NSCLC and post MPN AML *are being discussed with multiple institutions given the interest of the scientific community.*





Key Indications for JBI - 778



Disease Indications	Rationale	JBI – 778 Response
Non-Small cell lung cancer (NSCLC) with or without brain metastases	 EGFR (epidermal growth factor receptor) mutations are observed in 10 - 50% of non-small cell lung cancer patients EGFR mutations is almost double in patients with NSCLC with CNS metastases compared with patients without CNS metastases Splicing mutation or deletion (U2AF1, RBM10, etc) are common in NSCLC (8-10%) Non-responders to EGFR 3rd generation inhibitors have enriched splicing mutations Recently PRMT5 inhibitors have been shown to be effective in NSCLC patients who is resistant to Osimertinib (3rd Generation EGFR inhibitors) 	 PRMT5 is involved in splicing Mutations PRMT5 Inhibitors are sensitive to spliceosome mutant cell line both in <i>vitro and in vivo</i> JBI-778 is potent PRMT5 inhibitor having good plasma and brain exposure and has a potential to treat patients who are non- responders to the EGFR inhibitors with or without brain metastases
High Grade Glioma	 High-grade gliomas account for approximately 15 to 20 % of CNS tumours in children and adolescents Isocitrate dehydrogenase (IDH) mutant gliomas are the most common malignant primary brain tumors diagnosed in patients younger than 50, an important cause of morbidity and mortality PRMT5 inhibitors shown CR in IDH+ patients but faced tox issues impeding further development 	 JBI-778 has superior brain exposure and substrate competitive binding has established superior safety in preclinical setting JBI-778 has shown excellent results in pre-clinical in vivo model of glioma

Continue to invest



Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Ү-о-Ү	FY23	FY24	Ү-о-Ү
Revenue	0	0	0		4	0	(100%)
EBITDA	(10)	(5)	(7)	30%	(35)	(30)	14%

Continue to invest in two clinical stage programs

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Consolidated Reported Financials – Q4 FY24 & FY24



Total Income growth (YoY) along with EBITDA margin expansion (YoY)

Particulars (Rs. Cr.)	Q4'FY23	Q3'FY24	Q4'FY24	Y-o-Y	FY23	FY24	Y-o-Y
Revenue	1,678	1,677	1,759	5%	6,282	6,703	7%
Other Income	4	36	14		38	69	
Total Income	1,683	1,713	1,773	5%	6,320	6,772	7%
EBITDA	240	267	289	20%	827	994	20%
EBITDA Margin (%)	14.3%	15.6%	16.3%	200 bps	13.1%	14.7%	160 bps
Impairment of assets	171	0	0		171	0	
Exceptional Item	0	0	169		57	169	
РВТ	(86)	101	(54)	38%	28	171	513%
Normalised PBT	85	101	115	35%	256	339	33%
Normalised PBT Margin	5.1%	5.9%	6.5%		4.1%	5.0%	
Reported PAT	(101)	66	(62)	39%	(65)	73	212%
Normalised PAT ¹	27	66	61	122%	120	195	63%

FY24 Total Income grew YoY on the back of growth in revenue in Radiopharma, Allergy Immunotherapy and other income

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- FY24 EBITDA margin expanded YoY, led by overall improved margins in Radiopharma, Allergy Immunotherapy, Generics and API business
- FY24 normalised PAT grew YoY due to improved operating performance and increase in other income
- Exceptional Items for FY24 primarily includes Impairment of PPE and other intangible assets, pursuant to closure of manufacturing operations at solid dosage formulation facility at Salisbury, Maryland USA.

1. Normalised PBT / PAT is after adjusting for exceptional item & Impairment Charges

Key Ratios Net Debt / Ebitda is moving in the right direction

Particulars (Rs. Cr.)	Mar 31, 2023	Mar 31, 2024		
Gross Debt (Net of DIC)	3,410	3,414		
Net Debt (On constant currency)	2,426	2,464		
Net Debt / Ebitda	2.93	2.48		

- Strong focus on working capital and cash flow management
- Net cash generated from operating activities increased from Rs. 661 Cr. in FY23 to Rs. 971 Cr. in FY24

Sustainability



Signed PPA¹ & SHA² to purchase renewable energy for 92% of electricity demand by JPM entities in Karnataka in FY24



Summary – Q4'FY24 & FY24



1 Radio Pharmaceuticals : New products Mertiatide & Sulfur Colloid and Ruby-Fill® driving growth momentum Radio Pharmacies : Volume led growth & operational efficiencies driving margin expansion	
2 Allergy Immunotherapy : Sustained growth momentum & EBITDA margins	
3 CDMO Sterile Injectable : Capacity expansion at Spokane on track. Expect Line 3 to start commercial operations by Q1FY26	
4 CRDMO DDS: US Biotech industry headwinds continue. Medium term outlook continues to be positive CRDMO API : Pricing pressure continues. Taking initiatives to reduce operating costs & increase capacity utilization	

Generics : USFDA determines "VAI status" at Roorkee facility. Target to **reach EBITDA breakeven**

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Prop Novel Drugs : Preparing for Phase 2 Clinical trails and Investigator trials in JBI-802

Financial Results Table



Total Income (Rs. Cr.)	Q4'FY23		Q3'FY24		Q4'FY24		FY23		FY24	
Revenue (A)	1,678		1,677		1,759		6,282		6,703	
a. Radiopharma	689		752		818		2,552		3,001	
Radiopharmaceuticals	215		241		256		872		952	
Radiopharmacies	475		511		561		1,681		2,050	
b. Allergy Immunotherapy	170		161		188		603		679	
c. CDMO Sterile Injectables	321		303		259		1,155		1,117	
d. Generics	199		199		201		762		775	
e. CRDMO	294		252		282		1,185		1,093	
Drug Discovery Services	131		114		117		522		449	
CDMO – API	163		138		165		663		645	
f. Proprietary Novel Drugs	0		0		0		4		0	
Unallocable Corporate Income	5		11		11		22		38	
Other Income (B)	4		36		14		38		69	
Total Income (A+B)	1,683		1,713		1,773		6,320		6,772	
EBITDA (Rs. Cr.)	Q4'FY23	Margin	Q3'FY24	Margin	Q4'FY24	Margin	FY23	Margin	FY24	Margin
a. Radiopharma	112	16%	175	23%	169	21%	391	15%	584	19%
Radiopharmaceuticals	100	47%	126	52%	126	49%	465	53%	477	50%
Radiopharmacies	(4)	(1%)	10	2%	38	7%	(87)	(5%)	56	3%
b. Allergy Immunotherapy	55	33%	62	38%	75	40%	206	34%	273	40%
c. CDMO Sterile Injectables	86	27%	37	12%	58	22%	345	30%	192	17%
d. Generics	(39)	(20%)	(31)	(15%)	(39)	(19%)	(230)	(30%)	(141)	(18%)
e. CRDMO	47	16%	41	16%	52	19%	199	17%	169	15%
Drug Discovery Services	35	26%	30	27%	29	24%	164	31%	106	24%
CDMO – API	12	7%	11	8%	24	14%	35	5%	63	10%
f. Proprietary Novel Drugs	(10)		(5)		(7)		(35)		(30)	
Unallocable Corporate (Expenses) / Income	(11)		(13)		(19)		(49)		(55)	
Total EBITDA	240	14.3%	267	15.6%	289	16.3%	827	13.1%	994	14.7%

Note : "Radiopharma" segment EBITDA includes "EBITDA share" & "Share of profit" from Sofie

Annexure



JPM Business Strategy

To strengthen the unique position of each of the business unit to enhance shareholder value



INNOVATE

Radiopharma



- Continue to grow existing radiopharmaceutical products & launch new products
- Gain market share & increase profitability in radiopharmacies



2

Allergy Immunotherapy



- Gain share in the
 US Allergenic
 extracts
- Enlarge US Venom
 market
- Penetrate outside US markets



3

CDMO Sterile Injectables



- Double the capacity to leverage demand – supply gap in the finish space
- Leverage strong
 customer
 relationships to fill up
 the new capacity







Diversify the customer segments by tapping into large Pharma

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- Strengthen capabilities in development
- Leverage the partnership with Biotechnology companies



GENERICS

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- Non-US(International): Grow the business profitably
- India : Build 3 to 4 therapeutic areas in branded generics
- US : Make business profitable through change of operating model

DISCOVER

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PROPRIETARY NEW DRUGS



- All programs on track. Phase 1 data for JBI-802 Indicates therapeutic potential
- To explore institutional funding post early phase 2 data for JBI-802

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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







Dr. Tushar Gupta SVP – Corporate Strategy

Executive Leadership Team





Harsher Singh CEO - Jubilant Radiopharma



Giuliano Perfetti CEO - CRDMO, Biosys



Kyle Ferguson CEO – Allergy Business



Dr. Jaidev Rajpal CEO - Jubilant Generics



Chris Preti CEO - CDMO



Dr. Syed Kazmi CEO - Jubilant Therapeutics

Shareholding Pattern







GLOSSARY



Abbreviation	Details	Abbreviation	Details
CVS	Cardiovascular System	MEA	Middle East Africa
CNS	Central Nervous System	LATAM	Latin America
CDMO	Contract Development Manufacturing Organization	LOE	Loss of exclusivity
CRDMO	Contract Research & Development Manufacturing Organization	FDA (US)	U.S. Food and Drug Administration
F18	Fluorine-18 Radioisotope	PMDA (Japan)	Pharmaceutical and Medical Device Agency
PSMA	Prostate Specific Membrane Antigen	KEDA (Koroa)	Karaa Eaad Davalanmant Authority
Lu177	Lutetium-177 Radioisotope	AND/ICA (Roreal)	No Benetikas Haakka Dasa Astara
Ac225	Actinium-225 Radioisotope	ANVISA (Brazil) Brazilian Health Regulatory Agency
MAA	Macro Aggregated Albumin	TGA (Australia) Therapeutic Goods Administration
DTPA	Diethylenetriaminepentacetic Acid-Chelating Agent	API	Active Pharmaceutical Ingredient
HICON	Pharmaceutical Grade Radioactive Iodine	MENA	Middle East North Africa
I 131	Iodine-131 Radioisotope	GMP	Good Manufacturing Practices
MIBG	Metaiodobenzylguanidine	B2B2C	Business-to-Business-to-Consumer
USP (USP 825 Guideline)	U.S. Pharmacopeia (USP) general chapter ,825 (Related to Radiopharmaceuticals: Preparation,	B2B	Business-to-Business
Ca 69	Callium 68 Padiaisatana	ET/MPN	Essential thrombocythemia / Myeloproliferative neoplasm (rare chronic blood cancer)
Galos	Gaillum-68 Radioisotope	coREST	CRISPR-Cas9 Endomorphic RNA Symptomatic Inhibitor (RNA based therapy targeting genetic
RD	Rubialum (chemical element)	Epigenetic	disease)
Sr		Modulating	Medications that modify gene expression patterns
Cu 64	Copper-64 Radioisotope	Agent	
NRC	Nuclear Regulatory Commission (U.S.)	PRMT5	Protein Arginine Methyltransferase 5 inhibtor (Blocks enzyme activity involved in adding methy
GPOs	Group Purchasing Organisation	Brain	
IDNs	Integrated Delivery Network	Penetrant	Cerebral blood flow enhancers or cognitive-enhancing drugs (supplements)
SCIL	Sublingual immunotherapy (Allergy treatment - Dust mites & Seasonal allergy)	PD-L1 Inhibito	Programmed death Ligand-1 inhibitor (blocks the PD-L1 pathway, enhancing immune response
SCIT	Subcutaneous Immunotherapy (Allergy treatment Insect venom, pet dander, Mold, and other allergens)	PAD4 Inhibitor	poly(ADP-ribose) polymerase 4 inhibitor (Disrupts DNA repair mechanisms in cancer cells,
APAC	Asia Pacific	LSD1/HDAC6	Lysine specific demethylase 1/Histone deacetylase 6 inhibtor (Blocks enzymes involved in
MEA	Middle East Africa	inhibitor	modifying histones, impacting gene expression regulation in cancer therapy)
NSCLC	Non-small cell lung cancer	NSCLC	Non-small cell lung cancer
SCLC	Small cell lung cancer	SCLC	Small cell lung cancer

For More Information



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