

Bharat Parenterals Limited

Registered Office & Works: Survey No.: 144-A, Jarod-Samlaya Road, Vill. Haripura, Ta. Savli, Dist. Vadodara - 391520 (Guj.) India.

Mobile : 99099 28332

E-mail: info@bplindia.in, Web.: www.bplindia.in CIN NO: L24231GJ1992PLC018237

(WHO-GMP CERTIFIED ★ STAR EXPORT HOUSE)

Date: 28.07.2025

To,
Secretary
Listing Department
BSE Limited
Department of Corporate Services
Phiroze Jeejeebhoy Towers,
Dalal Street, Mumbai-400001.

Script Code: 541096

Dear Sir/Madam,

Subject: Investor Presentation for the quarter ended June 30, 2025.

Pursuant to Regulation 30 of the SEBI (Listing Obligations and Disclosure Requirements) Regulations, 2015, please find enclosed herewith the Investor Presentation for the quarter ended June 30, 2025.

Kindly take the same on your record. Thanking You,

Yours faithfully,

For, Bharat Parenterals Limited

Mr. Sharmin Soni Company Secretary & Compliance Officer

M.No: A-75694



Except for the historical information contained herein, statements in this presentation and the subsequent discussions, which include words or phrases such as "will", "aim", "will likely result", "would", "believe", "may", "expect", "will continue", "anticipate", "estimate", "intend", "plan", "contemplate", "seek to", "future", "objective", "goal", "likely", "project", "should", "potential", "will pursue" and similar expressions or variations of such expressions may constitute "forward-looking statements". These forward-looking statements involve a number of risks, uncertainties and other factors that could cause actual results to differ materially from those suggested by the forward-looking statements. These risks and uncertainties include, but are not limited to our ability to successfully implement our strategy, our growth and expansion plans, obtain regulatory approvals, our provisioning policies, technological changes, investment and business income, cash flow projections, our exposure to market risks as well as other risks. Bharat Parenterals Limited does not undertake any obligation to update forward-looking statements to reflect events or circumstances after the date thereof.

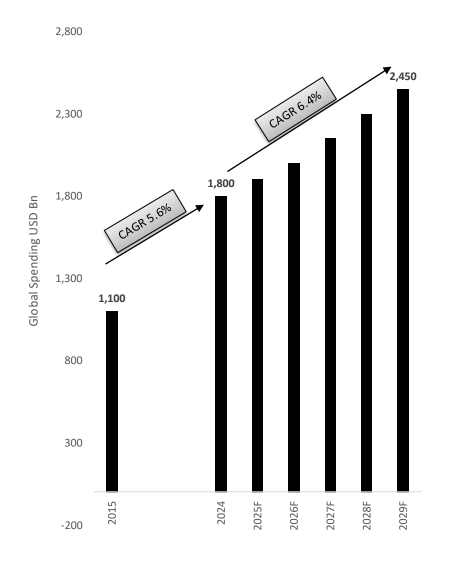
Table of contents

1	Market overview	4 - 7
2	Group overview	8 – 20
3	Financial update	21 - 25



Global medicine spending to reach \$ 2.4 trillion by 2029f with a few key themes having the greatest impact on growth and profitability

Global medicine market spending¹



Key themes in the generic finished dosage formulations space

1 GEOGRAPHY FOCUS

Higher growth and stable pricing in emerging markets

- High volume growth and negligible price erosion in emerging market generics vis-àvis regulated markets
- Evolving regulatory requirements have created entry barriers, reducing competition in emerging markets

2 NICHENESS OF PORTFOLIO

Superior margins and fewer competitors for niche portfolios

- Complex and specialty generics portfolios enjoy substantially higher margins across geographies
- Portfolios backed by innovative technology platforms have greater barriers to entry and fewer competitors

3 BRANDED GENERICS

> Strong brands enjoy stable market shares and pricing power

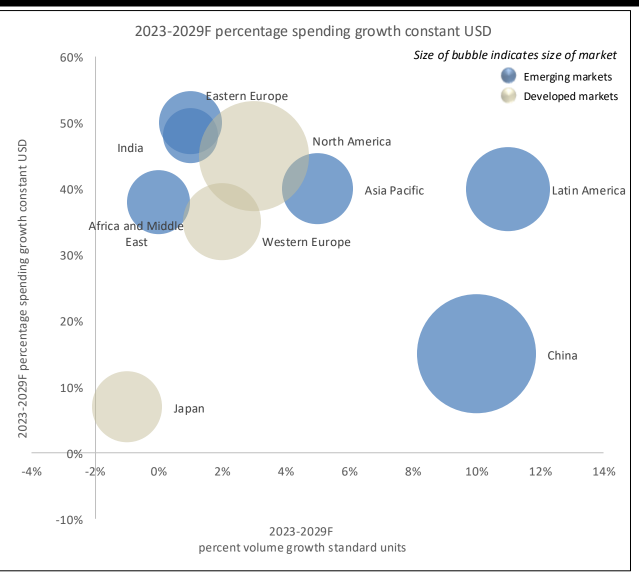
- Established brands command premium prices in emerging markets
- Once established, the market shares of top brands have remained stable over time

Source: IQVIA Market Prognosis, May 2025; IQVIA Institute, May 2025.

Note 1: Measures the amount spent purchasing medicines from manufacturers before off invoice discounts and rebates, and excludes the impact of spending on COVID 19 vaccines and therapeutics

Emerging markets expected to experience high growth in spending and volume, while both volume and spending growth to be muted in the developed markets

Population driven volumes and shift towards more expensive medicines because of improved healthcare penetration and rising per capital income will drive emerging market growth trends



Country	Growth trends	Volume and spending growth drivers
India LATAM	High volume growth	Population driven volume growth
APAC Africa & ME	High spending growth	 Spending growth from a shift in the product mix to more expensive products as healthcare access and per capita income levels improve
China	Moderate-high volume growth Muted spending growth	 Population driven volume growth Muted spending growth as more drugs are added to the NRDL and subjected to price negotiation
E. Europe	Low volume growth High spending growth	 Volume growth hampered by regional disruptions from Ukraine Spending driven by expected adoption of novel¹ drugs
W. Europe		Negligible volume growth – stagnant
N. America	Low volume growth	population/healthcare penetration growth
Japan	Low spending growth	 Spending growth driven by novel¹ drugs and offset by generic price erosion

Source: IQVIA Market Prognosis, May 2025; IQVIA Institute, may 2025.

LATAM: Latin America, E. Europe: Eastem Europe, APAC: Asia Pacific, ME: Middle East, W. Europe: Western Europe, N. America: North America, NRDL: National Reimbursement Drug List.

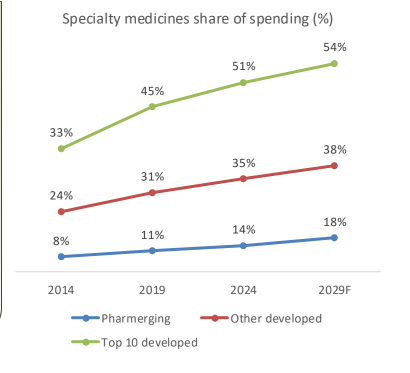
Note 1: Novel drugs are innovative drugs sold under the innovator brand

Branded generics in emerging markets and specialty medicines in developed markets expected to be the most rewarding spaces

2

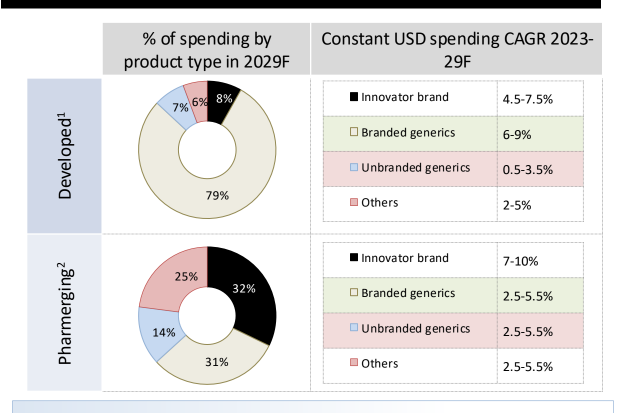
Specialty medicines will be one of the most rewarding spaces in developed markets as the share of spending on them continues to rise

- Specialty medicines are those which treat chronic, complex and rare diseases, and are characterized by complexity in storage, administration, distribution, and high prices
- Specialty medicines can be novel³ medicines or generics and are usually niche products



- In 2024, specialty medicines accounted for 51% of spending in the top 10 developed countries and 35% in other high and upper-middle income countries—up from 33% and 24% a decade ago.
- Specialty medicines make up 2–3% of volume but a growing share of spending. While they meet critical needs for few patients, costs for traditional therapies are declining.
- Pharmerging countries spent 14% on specialty medicines in 2024, projected to rise to 18% by 2029F, mainly limited by cost

The branded generics segment will be the most attractive in Pharmerging markets



- Wealthier countries spend more on original branded drugs, especially early in patent life
- Lower-income countries rely more on generics and branded generics (copy products).
- Pharmerging countries spend less on originators and more on low-cost generics or non-original brands

Source: IQVIA Market Prognosis, ; IQVIA Institute, May 2025



The BPL group is built to develop and manufacture FDFs for global markets...

Group overview

Bharat Parenterals Pvt. Ltd. (listed holding company)

Focus: Export-led pharmaceutical manufacturer of finished

dosage forms (FDFs).

Key therapies: Anti-infectives, anaesthesia, pain, CVS

Key dosage forms: Injectables, tablets, capsules, eye/ear drops

Key geographies: India, Africa, LATAM, SEA, ME

Standalone

REVENUE FY25: ₹318.7Cr EBITDA FY25: ₹48.1Cr PAT FY25: ₹26.4Cr

100% subsidiary

Capex FY25: ₹14+ Cr



Details
Vadodara, Gujarat
~28,500 sq. mt
~14,300 sq. mt
~4,300 sq. mt

100% subsidiary

Innoxel Lifesciences

55.9% subsidiary

Focus: Development and manufacturing of complex/specialty drugs for developed markets

Key Therapies: Oncology, pain management, Alzheimer's, long-acting injectibles and liquids

Pipeline portfolio overview: 40+ complex products (majority 505(b)(2) and

ANDAs); 10+ partnered with global clients

Key geographies: US (majority) and Western Europe

The company is driven by a well-balanced founding team, with 55.9% ownership by promoters and the remaining equity held by experienced technocrats.

Varenyam Healthcare

Focus: Branded generics for India's institutional market.

Key Therapies: Anesthesia, critical care, pain management. Expanding into complex general & oncology injectables (via Innoxel)

Key Geographies: Pan-India presence across major hospital chains.

Team & Strengths: 180+ on-ground reps across metros and Tier 1/2 cities.

Strong hospital-led channel, not retail/PCD focused.

Varenyam Bio Lifesciences

Focus: Manufacturing complex injectables/Specialty Drugs for regulated emerging markets

Acts as a complementary platform to Innoxel, extending global reach

Key Therapies: Complex generics across oncology, long-acting

Key Geographies: Emerging markets

injectables, NDDS – leveraging Innoxel's pipeline

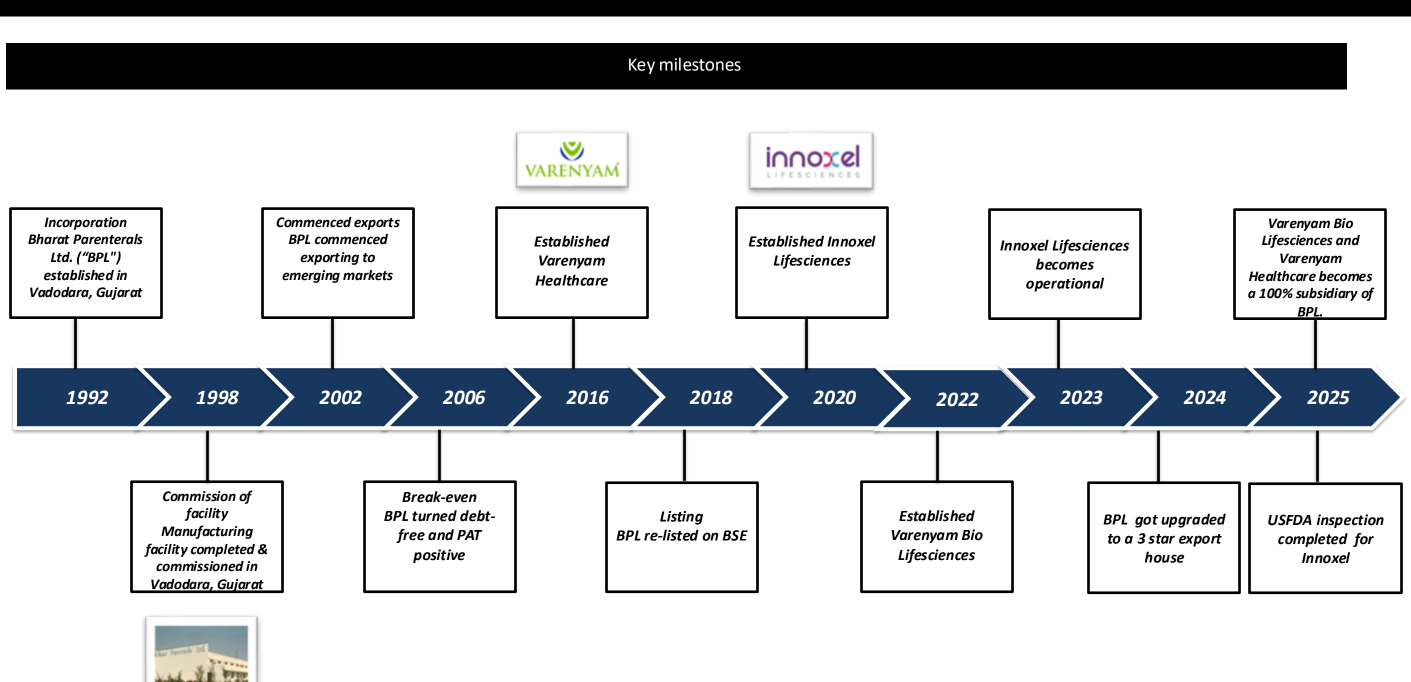
Pipeline & Strategy: Will use Innoxel's validated products under royalty-based arrangement. Reduces time-to-market by avoiding repeat development

Facility Status: Under construction; targeted operational readiness by FY27

Capex till date ~₹30Cr

Minimal Direct Capex

Capex ~₹250Cr Facility fully constructed, inspected by USFDA and undergoing product validation



...Poised to achieve rapid revenue growth and margin expansion over the next few years...

1 *Solid core business* primed for growth and margin expansion

(B) Bharat Parenterals Limited

Deep entrenchment in high-growth geographies enabled by experience of 3+ decades

Regularly upgraded manufacturing facility through the decade

Thoughtfully curated pipeline of product registrations designed to achieve revenue growth and realign product mix to yield higher margins

Promising pipeline driven by world-class R&D with the potential to create a durable, high-margin business



Founding team with the perfect blend of skills to create a regulated market CDMO success story

Supported by a truly state-of-the-art manufacturing infrastructure for the US and EU markets

Differentiated technology platforms with the potential to solve unmet healthcare needs, and a demonstrated track record of commercial success

Strengths across the CDMO continuum to address the complexities of the technology platforms

Promising pipeline that is highly market attuned and leverages the group's experience and expertise

volume and segment expansion.



Integration with BPL's manufacturing to drive

Leveraging complex product portfolio and market access for continued expansion

Strong presence in top Indian hospitals with proven execution in anesthesia and pain management; first in India to launch Sugammadex

Leveraging BPL's manufacturing and F&D capabilities to enter complex and niche markets within India.

Leveraging Innoxel's complex product portfolio and BPL's market access to achieve further expansion

Expansion into emerging regulated markets using Varenyam Bio infrastructure

Regulatorily accredited manufacturing infrastructure

Product registration pipeline aims to diversify geography mix First-time filings in new countries to New product filings in select existing Region expand presence within the geography countries to deepen presence LATAM Guatemala Ecuador Paraguay Nicaragua Peru Venezuela Honduras El Salvador APAC Uzbekistan Cambodia **Philippines** Kyrgyzstan Nepal Myanmar Vietnam Afghanistan Sri Lanka Africa and ME **Ivory Coast** Zambia Uganda Ethiopia Tanzania Nigeria Madagascar Ghana Mozambique Kenya Iraq Malawi EU Georgia Kosovo

Diversification to achieve growth and margin expansion

Growth objective

• Enhanced focus on APAC and LATAM that have higher volume and value growth vs. Africa

Margin expansion objective

ning

 Across regions, BPL is prioritizing countries where stringent regulatory and compliance requirements have created high entry barriers, resulting in fewer competitors and higher margins

Commissioning one new EU GMP compliant blocks

 Post approval these blocks will enable access to several APAC geographies that accept EU GMP compliant manufacturing facilities

Regulatory accreditations





















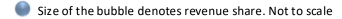






...Therapy area focus, and product mix

Therapy area	Sound strategy guiding therapy are	a-wise objective	Thoughtfully designed product registration pipeline		
	Strategy	Current	Future		
Anti-infectives	 Anti-infectives are competitive spaces with moderate margins BPL plans to shift focus away from anti-infectives into other categories Realign focus to select higher-margin products 	X		 BPL has selectively filed newer classes of antibiotics like Tigecycline, Tazobactam, and other niche anti-infectives Limited filing of older generation anti-infectives 	
Critical care	 Injectable products in this category have few competitors and higher margins BPL plans to expand presence and increase revenue contribution from this portfolio 		•	 Renewed focus on critical care products like Bupivacaine, Lidocaine, Atracurium Besylate with filings of these products in new geographies Filed higher-margin anaesthesia products like Sugammadex Filed higher-margin pain products like Tramadol and Pentazocine 	
Others	 Enter niche products with higher margins across a variety of therapeutic categories to replace anti- infectives 		•	Filed higher-margin products in CNS (Fluphenazine Decanoate) and CVS (Glyburide + Metformin)	



Innoxel at a glance

Innoxel is an innovation-driven, regulated market CDMO with their own product portfolio of specialty generics

Snapshot

- Overview:
- Innoxel is a regulated market focused, specialty FDF CDMO.
- ➤ Engaged in the development, manufacturing, and partnering of complex dosage forms, 505 (b)(2)s, and other specialty products
- Primary markets: US and Europe

Portfolio overview

Innoxel's portfolio provides solutions for unmet healthcare needs, and has been built around an identified set of differentiated technology platforms, which leverage the founding team's experience and expertise



- Capacity of 6 mn vials p.a.¹
- Expandable to 14 mn p.a. per line (General and potent lines)



- Capacity of 3 mn bottles p.a.²
- Expandable to 6 mn p.a. per line (General and potent lines)

Business segments

Portfolio of own products which have been outlicensed to front-end marketing partners for milestone payments + transfer revenues + profit share

CMO contracts with innovator and generic large pharma, yielding conversion-cost-based revenues

Capabilities and capacity

Oral liquid formulations



Liposomal injectables



Extended release injectables



Infrastructure



Located in Vadodara with a total manufacturing area of 350,000 saft



2 manufacturing blocks

Block 1 - General manufacturing Oral liquids in bottles and injectable vials Block 2 - Oncology manufacturing Oral liquids in bottles and injectable vials

Planned regulatory approvals



Successful completion of the USFDA inspection in May 25.



Regulatory inspection anticipated by Q3 FY26

- India's only US FDA approved Oncology Oral Liquid manufacturing facility, and one of only six globally.
- One of the few SKID based manufacturing facilities in India.

Business segment overview

Innoxel has two business segments with revenues from manufacturing, milestone achievements, and profit share from clients and partners

Segments		Own products	СМО			
Description	after which it is through to filinPartner owns t marketing.	es and carries out product development up to a certain stage, sout-licensed to a front-end partner who will fund the product g and approval. he NDA/ANDA/MA and will be responsible for front-end- the exclusive manufacturer for the product	 Innoxel manufactures the product for their client, providing manufacturing support from the clinical trial stage to the commercial manufacturing stage. One of the only India-based formulation CMOs working with Innovator clients for their novel molecule 			
	Manufacturing	Revenue (at an agreed upon transfer price) from contract manufacturing of products for front-end partner	Manufacturing	Revenue based on conversion cost per batch of manufacturing for outsourcing client		
Revenue streams	Milestones	Revenues tied to completion of clinical and product development milestones		Revenues tied to completion of clinical and product development milestones		
	Profit share	Pre-determined share of front-end partner's profits after accounting for transfer cost and marketing costs				
No of products	more are in vario	ned partnerships for 12 products with a front-end partner, while 20 us stages of development. Going forward, we plan to add 5–6 ar to achieve a diversified portfolio of 40+ products.	10 CMO contracts identified and signed. Several others in pipeline			
Client type		specialty generic companies with strong front-end presence in the ack record of successfully marketing specialty products		novator pharma companies requiring regulatorily approved city for complex products		

Pillars of a regulated market CDMO success story

Operational excellence

Mr. Bharat Desai



30+ years at Holdco managing a large injectable manufacturing company

Work experience:





 B.Sc (Chemistry) from SP University Differentiated R&D skills

Dr. Manish Umrethia



CEO of Auxilia Pharma, an R&D and formulation development company

Work experience:







- B.Pharm, M.Pharm (LMCP, Ahmedabad)
- Ph.D. (MS University of Baroda)
- Post Doctoral (Queens University, Belfast)

Sound strategic direction

Mr. Bhahim Desai



Managing Director of Varenyam Healthcare Pvt. Ltd, a domestic branded formulations company

Work experience:





- B.Pharm
- MBA in Pharmaceutical Marketing and Management, NMIMS, Mumbai

Wide clinical experience

Mr. Manoj Vyas



CEO of CBCC Global Research, a Contract Research Organisation based out of US and India

Work experience:



- M.Sc. Chemistry (Gujarat University)
- Masters Clinical Research (Cranfield University, UK)

Robust regulatory & compliance

Mr. Tushar Patel



CEO of Pharmazone, a provider of regulatory affairs and compliance advisory services

Work experience:



- B.Pharm. (LMCP, Ahmedabad)
- Masters Clinical Research (Cranfield University, UK)

Deep commercial networks

Mr. Manoj Bharathi



Director of GeneriQ Pharmaceuticals, a commercial licensing advisory firm

Work experience:



- B.Tech .Chemical Engineering (Anna University, Chennai)
- MBA (IIFT, Delhi)

Drug characteristics		Impact	
The encapsulated drug is protected from rapid degradation and elimination by the body	Ø		
The drug circulates in the body for longer, allowing for modified drug release profiles (sustained/controlled)	Ø		
Usually manufactured with naturally derived starting materials. Offer excellent biocompatibility and safety and fewer side effects		6	
Allow for targeted delivery of drug to site of disease and improved bioavailability. This improves therapeutic benefits and causes fewer side effects	Ø	6	
Well-suited for oncology			
Liposomal injectables are lipid-based drug vesicles with one or more bilayers enclosing an aqueous compartment.			
They can carry a hydrophilic drug in the aqueous compartment and a hydrophobic drug between the bilayers			
Lower dosage frequency which reduces discomfort and enhances patience convenience			
Ability to target specific anatomical sites in the body where high drug concentrations can be maintained. This improves therapeutic benefits and causes fewer side effects	Ø	6	
Improved patient compliance	Ø		
Allows for consistent levels of drugs in the body - fewer side effects and improved therapeutic benefits	Ø	6	
Well-suited for CNS disorders, chronic pain, hormonal contraception, and oncology			
• Extended release injectables are parenteral, sustained drug delivery systems which are injected into the body and then slowly released over a long period of time (typically 2-12 weeks)			
Oral liquids are absorbed more quickly compared to oral solids	Ø		
Convenience and comfort to pediatric and geriatric populations that struggle with swallowing solid orals		6	
Offer dosing flexibility. Simple and convenient to change the dosage in case of medicines requiring complex dose titration/adjustment based on body weight	Ø		
Well-suited for anti-hypertensives and CNS disorders			
Ready to use injectables ("RTU")			
Products with clinical complexity requiring patient based clinical trials (usually, generic product trials are carried out on healthy patients).			
Formulations with APIs that are difficult to source			
	 The encapsulated drug is protected from rapid degradation and elimination by the body The drug circulates in the body for longer, allowing for modified drug release profiles (sustained/controlled) Usually manufactured with naturally derived starting materials. Offer excellent blocompatibility and safety and fewer side effects Allow for targeted delivery of drug to site of disease and improved bioavailability. This improves therapeutic benefits and causes fewer side effects Well-suited for oncology Uposomal injectables are lipid-based drug vesicles with one or more bilayers enclosing an aqueous compartment. They can carry a hydrophilic drug in the aqueous compartment and a hydrophobic drug between the bilayers Lower dosage frequency which reduces discomfort and enhances patience convenience Ability to target specific anatomical sites in the body where high drug concentrations can be maintained. This improves therapeutic benefits and causes fewer side effects Improved patient compliance Allows for consistent levels of drugs in the body - fewer side effects and improved therapeutic benefits Well-suited for CNS disorders, chronic pain, hormonal contraception, and oncology Extended release injectables are parenteral, sustained drug delivery systems which are injected into the body and then slowly released over a long period of time (typically 2-12 weeks) Oral liquids are absorbed more quickly compared to oral solids Convenience and comfort to pediatric and geriatric populations that struggle with swallowing solid orals Offer dosing flexibility. Simple and convenient to change the dosage in case of medicines requiring complex dose titration/adjustment based on body weight Well-suited for anti-hypertensives and CNS disorders Ready to use injectables ("RTU") Products with clin	The encapsulated drug is protected from rapid degradation and elimination by the body The drug circulates in the body for longer, allowing for modified drug release profiles (sustained/controlled) Usually manufactured with naturally derived starting materials. Offer excellent biocompatibility and safety and fewer side effects Allow for targeted delivery of drug to site of disease and improved bioavaliability. This improves therapeutic benefits and causes fewer side effects Well-suited for oncology Liposomal injectables are lipid-based drug vesicles with one or more bilayers enclosing an aqueous compartment. They can carry a hydrophilic drug in the aqueous compartment and a hydrophobic drug between the bilayers Lower dosage frequency which reduces disconfort and enhances patience convenience Ability to target specific anatomical sizes in the body where high drug concentrations can be maintained. This improves therapeutic benefits and causes fewer side effects Improved patient compliance Allows for consistent levels of drugs in the body - fewer side effects and improved therapeutic benefits Well-suited for CNS disorders, chronic pain, hormonal contraception, and oncology Extended release injectables are parenteral, sustained drug delivery systems which are injected into the body and then slowly released over a long period of time (typically 2-12 weeks) Oral liquids are absorbed more quickly compared to oral solids Convenience and comfort to pediatric and geratric populations that struggle with swallowing solid orals Offer dosing flexibility. Simple and convenient to change the dosage in case of medicines requiring complex dose titration/adjustment based on body weight Well-suited for anti-hypertensives and CNS disorders Products with clinical complexity requiring patient based clinical trials (usually, generic product trials are carried out on healthy patients).	The encapsulated drug is protected from rapid degradation and elimination by the body The drug circulates in the body for longer, allowing for modified drug release profiles (sustained/controlled) Usually manufactured with naturally derived starting materials. Offer excellent biocompatibility and safety and fewer side effects Allow for targeted delivery of drug to site of disease and improved bioavailability. This improves therapeutic benefits and causes fewer side effects Well-suited for oncology Usually manufactured with naturally derived starting materials. Offer excellent biocompatibility and safety and fewer side effects Well-suited for oncology Usually manufactured with naturally derived starting materials. Offer excellent biocompatibility and safety and fewer side effects Well-suited for oncology Usually manufactured with naturally derived starting materials. Offer excellent biocompatibility and safety and causes fewer side effects Well-suited for oncology Lower dosage frequency which reduces discomfort and enhances patience convenience Ability to target specific anatomical sizes in the body where high drug concentrations can be maintained. This improves therapeutic benefits and causes fewer side effects Well-suited for CNS disorders, chronic pain, hormonal contraception, and oncology Extended release injectables are parenteral, sustained drug delivery systems which are injected into the body and then slowly released over a long period of time (typically 2-12 weeks) Convenience and comfort to pediatric and geriatric populations that struggle with swallowing solid orals Offer dosing flex bility. Simple and convenient to change the dosage in case of medicines requiring complex dose titration/adjustment based on body weight Well-suited for anti-hypertensives and CNS disorders Ready to use injectables ("RTU") Products with clinical complexity requiring patient based clinical trials (usually, generic product trials are carried out on healthy patients).

...With the ability to formulate solutions for unmet healthcare needs...

Well-hedged against all types of risk through leverage of strengths across the CDMO continuum

Risks	Description	Innoxel's hedge
Development risk	 Inability to successfully complete formulation development/achieve clinical objectives in a timely manner 	 Dr. Manish has led the development of over half the currently marketed liposomal injectables Dr. Manish has 50 patents to his name as a lead scientist and 40+ formulations in developed and developing market.
Clinical trial risk	 Delay in obtaining slots with a clinical trial services provider, patient recruitment, formulation of study design and protocol Risk of cost and time overruns 	 The waiting period for clinical trial slots for Innoxel will be lower by 8-10 months due to its affiliation with CBCC, helping them avoid delays and cost overruns
Filing and approval risk	 Inability to make complete filings, delays in approval resulting from issues in communication 	 Mr. Tushar and CBCC's combined expertise and experience in managing regulatory affairs and FDA communications Dr. Manish's experience with filing similar products
Commercialization risk	Inability to generate demand and win market share	 The portfolio has been curated to ensure that it caters to clear unmet patient needs Mr. Manoj's experience with finding the right licensing partners, who have the access and expertise necessary to commercialize the product and win market share
Infrastructure/ Regulatory risk	Receipt of adverse feedback by regulatory authorities post facility audit	 Innoxel to leverage the experience of Mr Tushar, who is a seasoned GMP consulting professional Operational aspects of the company to be overseen by the Holdco leadership team



Varenyam Healthcare is a high-impact speciality branded generics business.

Overview

Established in 2016, Varenyam Healthcare is a specialty pharmaceutical company focused on critical care, anesthesia, and pain management.

Presence in 7,500+ hospitals across India, supported by a 180+ person field force.

Strong presence in top institutional chains including Apollo, NH, Fortis, Manipal to name a few.

Products aligned with BPL's manufacturing, enabling better control over quality, speed, and margin.

Products

Portfolio includes high-quality injectables tailored for hospital-driven therapies.

First in India to launch Sugammadex 100 mg/ml (anaesthesia reversal) in JV with BDR Pharma.

FoQas – time–temperature indicator to ensure cold-chain compliance for sensitive products.

Pipeline includes complex general and oncology dosage forms in upcoming launches.



Strategy & Differentiators

Deep institutional presence with focused therapeutic strategy and a skilled sales force.

Growth via expanding portfolio and tapping into complex formulations using BPL/Innoxel's R&D.

Launching two new therapeutic divisions over next 2–3 years.

Plans to scale revenue to ₹100 Cr by FY28.

PRODUCT PORTFOLIO

Leverages Innoxel's & BPL's R&D pipeline through inlicensing, reducing time-to-market and avoiding repeat development.

Focused on complex dosage forms (injectables, long-acting formulations, oncology, NDDS, Lyophilized injectables).

Designed to repurpose and relicense Innoxel's 505(b)(2) and complex ANDA portfolio for high-growth, underpenetrated markets.

Varenyam Bio

Lifesciences

Overview

Incorporated in 2022, Varenyam Bio is a strategic extension of the BPL group focused on manufacturing complex injectables and oral liquids for regulated emerging markets.

Created to complement Innoxel Lifesciences by serving countries outside the US/EU, including those requiring higher regulatory approvals.

Facility will target EUGMP and local regulatory approvals across LATAM, Africa, Eastern Europe, and Australia.

Strategy & Infrastructure

Offers rapid entry into Tier 2 global markets through localized regulatory strategies.

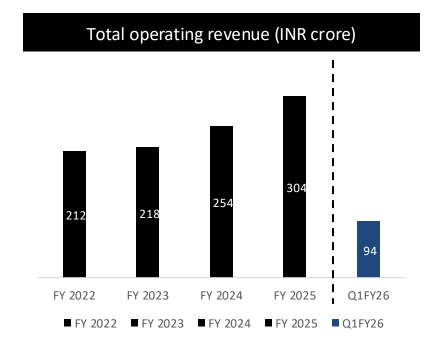
Lower-cost execution model with faster monetization than highly regulated CDMO pathways.

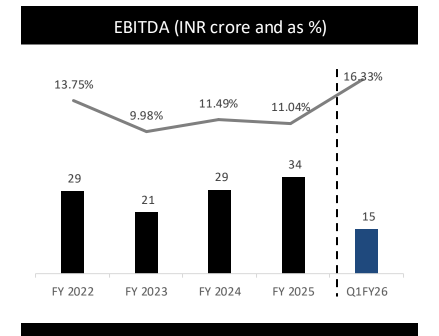
2 Particulate injectable lines: general and potent

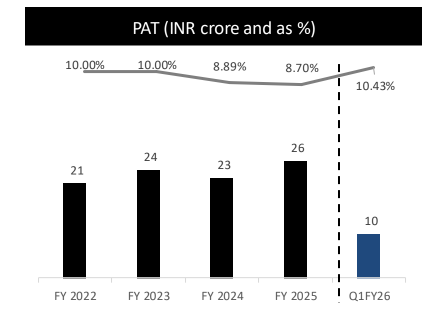
2 Lyophilized injectable lines: general and potent

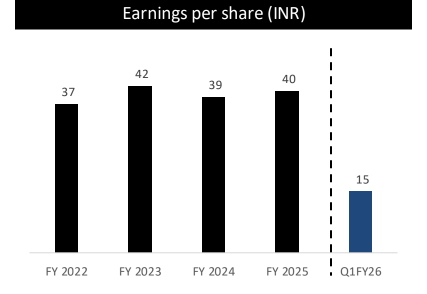


Financial metrics | Standalone







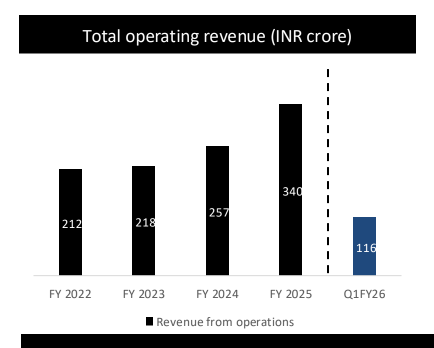


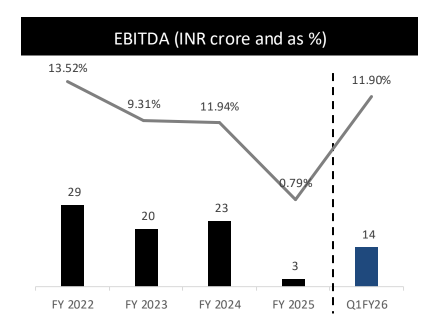
Financial metrics | Standalone key financials Q1 FY 26

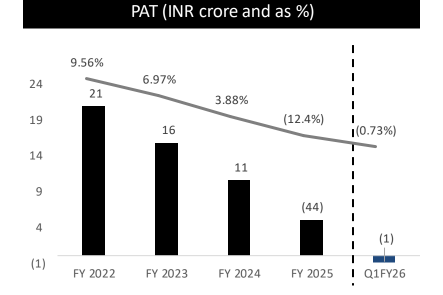
Figures in INR crore

Particulars	Q1 FY 2026	Q1 FY 2025	Change (%)	FY 2025	FY 2024	Change (%)
Revenue from operations	94.37	92.06	+2.51%	304.13	257.98	+17.89%
Other operating revenue	1.47	3.90		14.55	8.04	
Total operating revenue	95.84	95.96	-0.13%	318.68	266.02	+19.80%
EBITDA	15.41	14.99	+2.80%	33.59	29.21	+14.99%
EBITDA margin (%)	16.33%	16.28%		11.04%	11.49%	
PAT	10.00	12.18	-17.90%	26.44	22.59	+17.09%
PAT (%)	10.43%	12.69%		8.70%	8.89%	
EPS (INR)	15.26	20.26		40.36	38.97	

Financial metrics | Consolidated







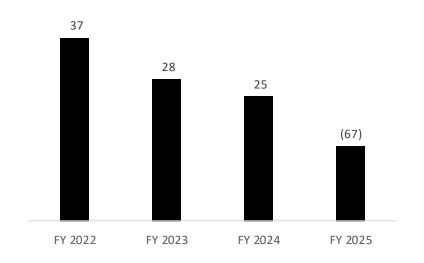
FY26E Financial Outlook

BPL (Standalone): The Company remains confident about its growth trajectory, maintaining FY26 revenue guidance of 12–14%, supported by a strong order book, expanded distribution footprint, and better capacity utilisation. EBITDA margins are expected to improve to 15–17%, driven by operating leverage and an increased share of institutional business.

Innoxel: Innoxel is expected to generate ₹65–70 crore in revenue in FY26, with a stable quarterly cost base of ₹15–16 crore. The business is poised to achieve operational breakeven through milestone-driven product revenues. Commercial CMO supplies are anticipated to commence in Q1 FY27, following the scheduled EU-GMP inspection in FY26.

Varenyam Healthcare: Revenue for Varenyam is projected at ₹60–65 crore in FY26, marking a 20–21% year-on-year growth. Expansion into new therapeutic areas and increased penetration in existing segments will continue to fuel momentum in the domestic business.

Earnings per share (INR)



Financial metrics | Consolidated key financials Q1 FY 26

Figures in INR crore

Particulars	Q1 FY 2026	Q1 FY 2025	Change (%)	FY 2025	FY 2024	Change (%)
Revenue from operations	116.00	92.51	+25.4%	340.38	261.22	+30.3%
Other operating revenue	1.06	2.54		11.90	4.68	
Total operating revenue	117.06	95.05	+23.2%	352.28	265.90	+32.5%
EBITDA	13.78	4.49	+206.9%	2.70	23.30	-88.4%
EBITDA margin (%)	11.9%	4.9%		0.79%	9.05%	
PAT	-0.86	-8.85	-90.3%	-43.67	10.51	-515.5%
PAT (%)	-0.73%	-9.31%		-12.40%	3.95%	
EPS (INR)	-1.27	-14.7		-66.64	24.94	