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May 10, 2024

To BSE Limited Phiroze Jeejeebhoy Towers, 25th Floor, Dalal Street, Mumbai - 400 001

The National Stock Exchange of India Ltd Exchange Plaza, Bandra Kurla Complex Bandra (E), Mumbai - 400 001

Scrip Code: 524558

Scrip Code: NEULANDLAB; Series: EQ

Dear Sir/Madam,

Sub: Investors/Analysts Presentation

Pursuant to Regulation 30 of the SEBI (Listing Obligations and Disclosure Requirements) Regulations, 2015, we are enclosing the presentation to the Investors/ Analysts on the Financial Results of the Company for the quarter and year ended March 31, 2024.

The presentation is also being uploaded on the website of the Company at www.neulandlabs.com

This is for your information and records.

Thanking you,

Yours Sincerely,

For Neuland Laboratories Limited

Sarada Bhamidipati Company Secretary

Encl: As above



Neuland Laboratories Limited

INVESTOR PRESENTATION Q4FY24 & FY24



SAFE HARBOUR

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 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. The Company does not undertake any obligation to update forward-looking statements to reflect events or circumstances after the date thereof.

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Q4FY24 & FY24 Highlights





SUCHETH DAVULURI

"We surpassed revenues of Rs. 1500 crores in FY24 with EBITDA at over 30%. This has been driven by high growth in the CMS business and steady growth of the Specialty GDS business, both of which were in line with our plans and expectations. Another key element is the work on optimizing costs & processes which will also make us truly sustainable."

SAHARSH DAVULURI

"Our CMS business saw robust growth in FY24 as some projects are near launch while key commercial products continue to scale. Our growing reputation and the macro-environment are ensuring that exciting opportunities come our way even as we work towards building a further differentiated customer experience. We will continue to invest for the future by adding capacity and capabilities."



Business and Financial Highlights





FY24 Business and Financial Highlights

CMS

CMS revenues driven by growth from commercial molecules and molecules close to commercialization.

GDS

Specialty business driven by Paliperidone and Dorzolamide In Prime segment Mirtazapine, Escitalopram and Levetiracetam were the key molecules

Regulatory Audits

US FDA inspected Unit-3 and issued EIR (Establishment Inspection Report) Unit-I inspected by EDQM (European Directorate for the Quality of Medicines) and the US FDA

Sustainability rating

S&P ESG rating of 64



Free Cash Flow (FCF) generation and utilisation

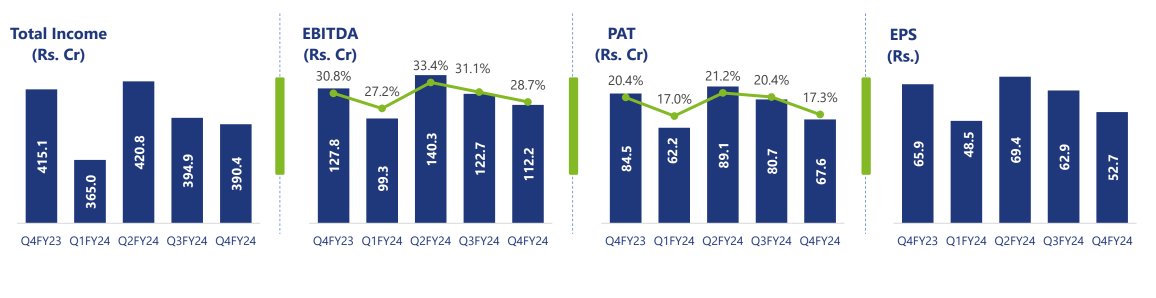
Generated Free Cash Flow of Rs. 116.4 crores during FY24, partly utilised to reduce debt by Rs 39.4 crores Capex Investment of Rs 143.7 crores for enhancement of capabilities

Working Capital

Reduction in working capital cycle to 122 days in FY24 as compared to 141 days in FY23

Q4FY24 Financial Highlights



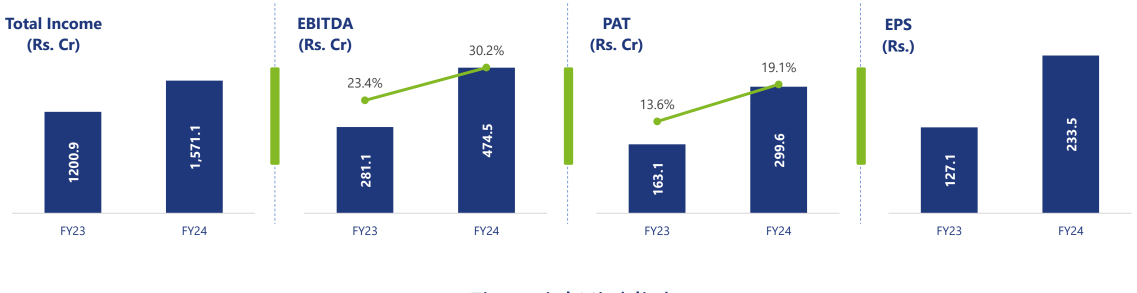




- Total Income for Q4FY24 at Rs. 390.4 crore (-6.0% YoY)
- EBITDA for Q4FY24 at Rs. 112.2 crore (-12.1% YoY)
- EBITDA Margin for Q4FY24 at 28.7% (decreased by 202 bps YoY)
- PAT for Q4FY24 at Rs. 67.6 crore (-20.0% YoY)
- Net Debt stood at Rs. (32.6) crore as at Q4FY24 end compared to Rs. 63.0 crore as at Q4FY23 end and Rs (44.6) crore as at Q3FY24 end

FY24 Financial Highlights

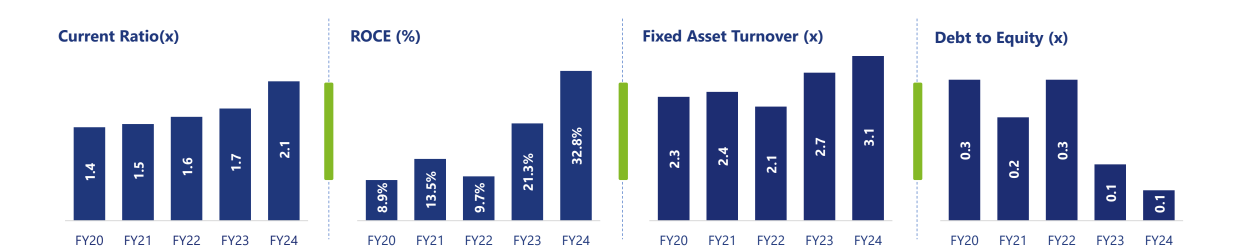






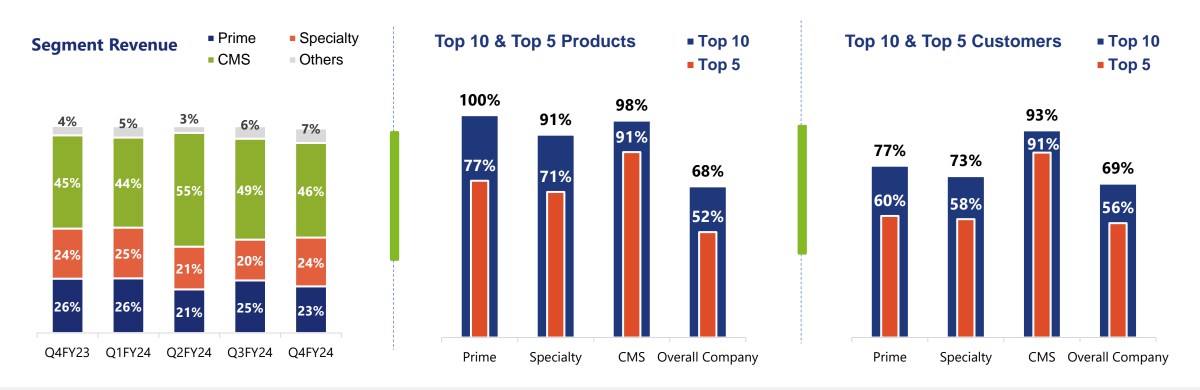
- Total Income for FY24 at Rs. 1,571.1 crore (+30.8% YoY)
- EBITDA for FY24 at Rs. 474.5 crore (+68.8% YoY)
- EBITDA Margin for FY24 at 30.2% (increased by 680 bps YoY)
- PAT for FY24 at Rs. 299.6 crore (+83.7% YoY)
- Net Debt stood at Rs. (32.6) crore as at FY24 end compared to Rs. 63.0 crore as at FY23 end





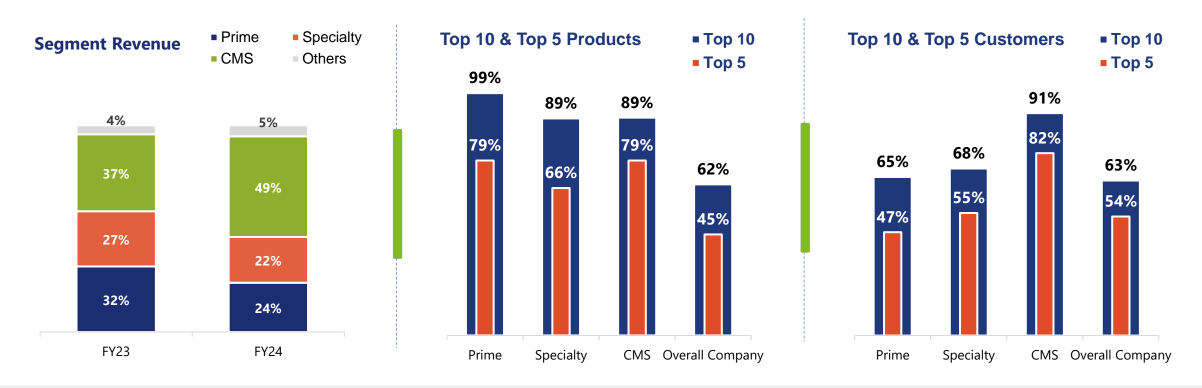
Particulars (Rs Cr)	Mar-20	Mar-21	Mar-22	Mar-23	Mar-24
Shareholder's Funds	705.5	781.9	835.6	988.4	1,276.5
Net Debt	199.9	152.1	212.0	63.0	-32.6
Tangible Assets (including CWIP and Investment property)	391.1	437.9	497.2	511.2	575.4
Working Capital	289.4	308.6	376.9	463.0	525.4





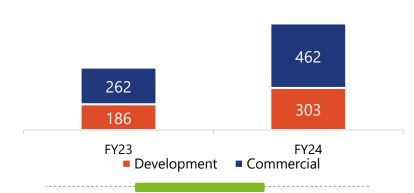
- Steady shift from low margin Prime to high margin Specialty and CMS segments
- CMS business caters to Innovator customers on an exclusive basis, developing and manufacturing APIs/Intermediates in line with rigorous customer expectations hence is highly concentrated in terms of customers
- Specialty segment works on complex products and technologies, hence has a focused approach towards select customers





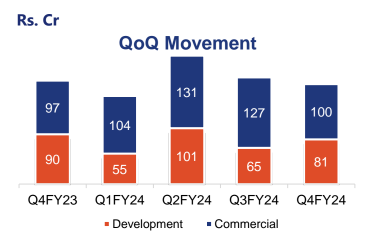
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YoY Movement

Rs. Cr



Q4FY24	Pre-Clinical	P-1	P-2	P-3	Pre-Reg/Reg*	Commercial	Total
API	8	8	12	3	8	8	47
Intermediate	8	4	9	4	6	10	41
Grand Total	16	12	21	7	14	18	88
Q3FY23	Pre-Clinical	P-1	P-2	P-3	Pre-Reg/Reg	Commercial	Total
API	15	5	7	4	8	9	48
Intermediate	10	4	4	2	7	12	39
Grand Total	25	9	11	6	15	21	87
Q3FY22	Pre-Clinical	P-1	P-2	P-3	Pre-Reg/Reg	Commercial	Total
API	14	3	7	7	8	8	47
Intermediate	7	5	2	0	8	12	34
Grand Total	21	8	9	7	16	20	81
Q3FY21	Pre-Clinical	P-1	P-2	P-3	Pre-Reg/Reg	Commercial	Total
API	15	3	7	3	12	6	46
Intermediate	7	4	2	0	8	11	32
Grand Total	22	7	9	3	20	17	78

No. of active CMS projects

• Pre-clinical to P-3: Neuland generates revenue by process research & development as well manufacturing quantities for clinical trials

- *Pre-Reg/Reg: Phase-3 complete; Molecules filed but not yet commercial (Earlier labelled as 'Development') or where customer working towards adding Neuland as a second source for a commercial molecule
- Commercial: Neuland generates revenues by manufacturing APIs for commercial novel molecules for innovators
- Steady trend in molecules transitioning from clinical phases to commercialisation resulting in increase in revenue from commercial products



Company overview

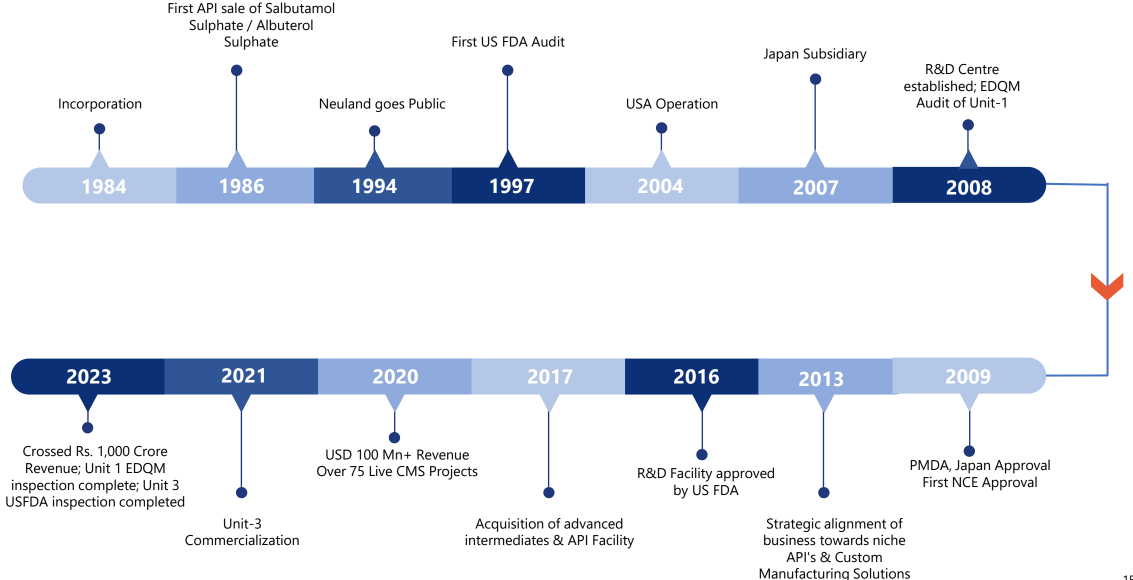


Key Facts



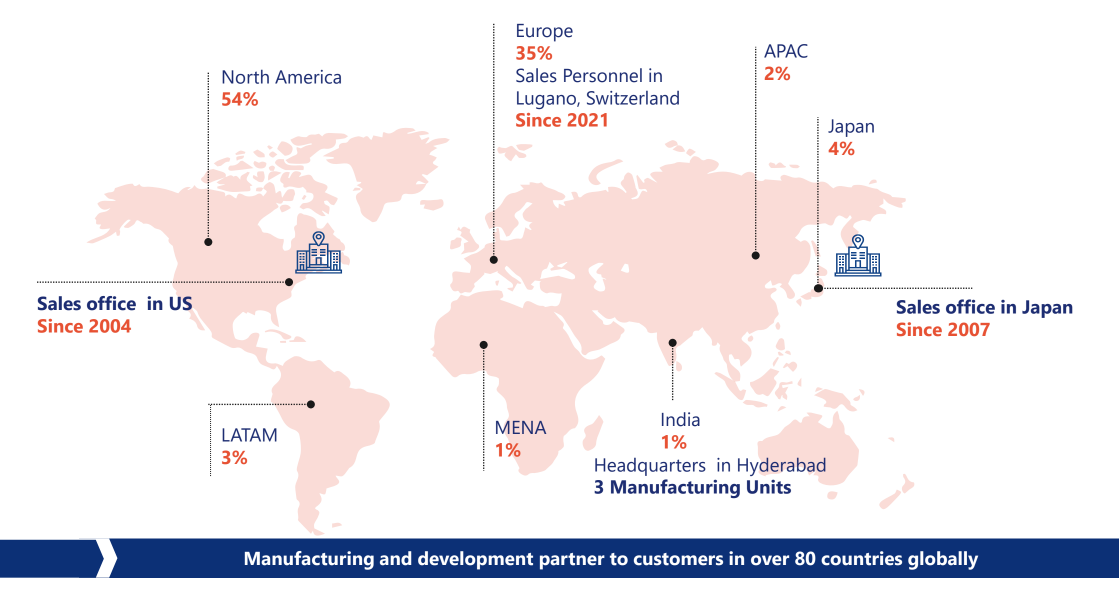






Our Reach





% refers to 9MFY24 Sales by End Market

Manufacturing Capabilities



	Established	Hydrogenation Reaction Volume	Solvent Recovery System	Cryogenic Reaction Volume	Regulatory	Total Reactor Volume
	1986	7.4KL	100KLD	25KL	USFDA, EDQM, CFDA, PMDA, et. al	233 KL
Unit-1 Unit-2	1994	6 KL	20KLD	15 KL	USFDA, EDQM, PMDA, ANVISA et. al	363 KL
Unit-3	2017 (Acquired)	Facility creation under process	50KLD	15KL	US FDA, EDQM, PMDA, ANVISA, et al.	305 KL

Priorities

- Modernizing and automation of overall operations
- Focus on adding capabilities and capacities
- Maintaining the leadership position in key molecules

Focused R&D Framework



Infrastructure

- ▶ 15 Development Labs with space for expansion
- ▶ 60 Fume hoods
- Analytical Labs
- Dedicated kilo Lab for Scale up
- Dedicated Labs for Peptides
- Separate facility for D2 analogues
- Approvals for Department of Scientific and Industrial Research (DSIR), Government of India and US FDA
- ▶ R&D team of 345 People



USFDA inspected Neuland's R&D facility in February 2016 without any observations

Significant R&D Achievements

- Several NCE APIs added in NDA or commercial stage drugs
- Support for multiple APIs each year in Phase 2 and Phase 3 clinical candidates
- ► Generic API business -
- 950+ DMFs filed
- 300+ API processes developed
- 204+ patents filed. Received USPTO patent for improved process synthesis of Paliperidone Palmitate
- ► 3 new DMFs filed in FY23

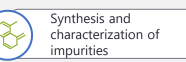
Analytical Capabilities



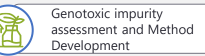
complex molecules



Complete analytical validation package (as per ICH guidelines)



Reference standard qualification



Study of Solid-state properties



Salt screening and optimization

Priorities

- Focus on quality enhancement and training for enhancement of technical skills
- Emphasis on complex molecules involving advanced chemistry, automation, upgradation of testing equipment, and complementary new technologies
- Consistent investments in Quality by Design (QBD) labs and process engineering







Rs. Cr

	FY2015	FY2016	FY2017	FY2018	FY2019	FY2020	FY2021	FY2022	FY2023	FY2024
Total Income	469.9	519.3	582.3	533.7	670.3	766.6	953.0	953.2	1,200.9	1,571.1
EBITDA	67.4	89.7	97.6	56.3	61.4	105.3	162.5	144.3	281.1	474.5
EBITDA Margin	14.3%	17.3%	16.8%	10.6%	9.2%	13.7%	17.1%	15.1%	23.4%	30.2%
PAT	15.9	34.8	41.4	13.6	16.1	15.9	80.3	63.5	163.1	299.6
PAT Margin	3.4%	6.7%	7.1%	2.5%	2.4%	2.1%	8.4%	6.7%	13.6%	19.1%
EPS	18.5	30.8	36.9	10.6	12.8	12.4	62.6	49.5	127.1	233.5
Current Ratio (x)	1.1	1.2	1.2	1.2	1.4	1.4	1.5	1.6	1.7	2.1
ROCE (%)	15.7%	18.4%	15.9%	5.0%	4.7%	8.9%	13.5%	9.7%	21.3%	32.8%
Fixed Asset Turnover (x)	3.8	3.7	3.8	3.2	2.9	2.3	2.4	2.1	2.7	3.1
Debt to Equity (x)	1.1	0.9	0.9	0.5	0.3	0.3	0.2	0.3	0.1	0.1

- Revenue was impacted in FY2018 as a result of mismatch in capacity vs orders. EBITDA margins in FY19 & FY20 were impacted as a result of spike in RM prices, which led Neuland to actively work towards Supply chain de-risking before the COVID19 pandemic
- ROCE was impacted by due to acquisition of unit III in FY2018 which was commercialized in FY2021. Unit 3 utilisation levels have recently started ramping up and momentum is expected to continue



Business 3 Strategy



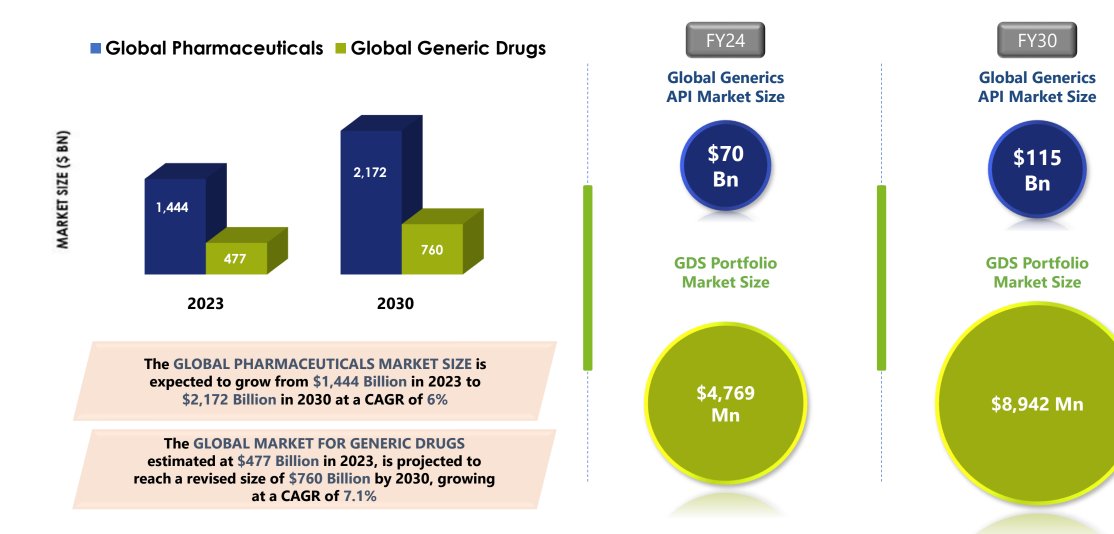




GDS Strategy

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1. Maximizing Current Portfolio

- Increase wallet share from existing customers
- Focus on regulated markets/ quality conscious customers
- Early identification for primary sourcing opportunities
- Exploring the additional opportunities from Line extension in terms of new dosage forms and indications
- Focus on customers with backward integration to convert them into alternate sourcing opportunities



- Commercialization of pipeline molecules through New leads identification and conversion
- Aim for first sourcing and NCE-1 opportunities
- Investment in new areas

••••

GDS

- Explore collaboration opportunities with dossier development companies
- Filing DMFs for peptides





CMS Strategy

Global Outlook of CDMO Industry



CDMOs are strengthening their position as indispensable partners and creating strategic, integrated partnerships with innovators

CDMOs are becoming more agile to meet increasing demand for diverse projects

> Rise in M&A and PE investments has resulted in increased focus on CDMO business

CDMO market expected to be driven by rising rates of chronic and lifestyle diseases

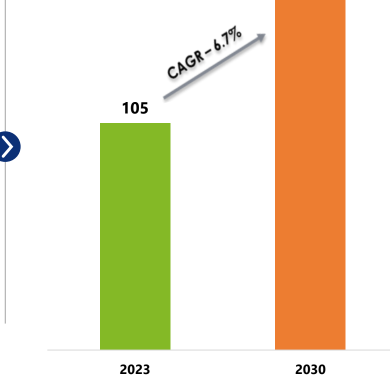
CDMOs are expanding beyond traditional technologies and producing advanced therapies while becoming end to end service providers



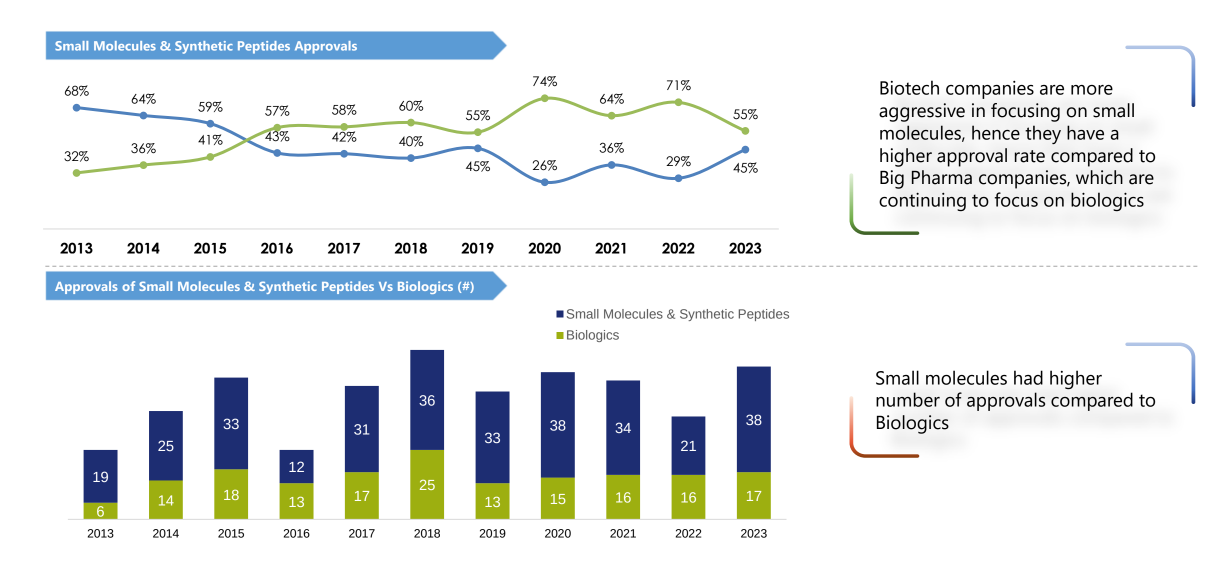
Big Pharma also acting as CDMOs



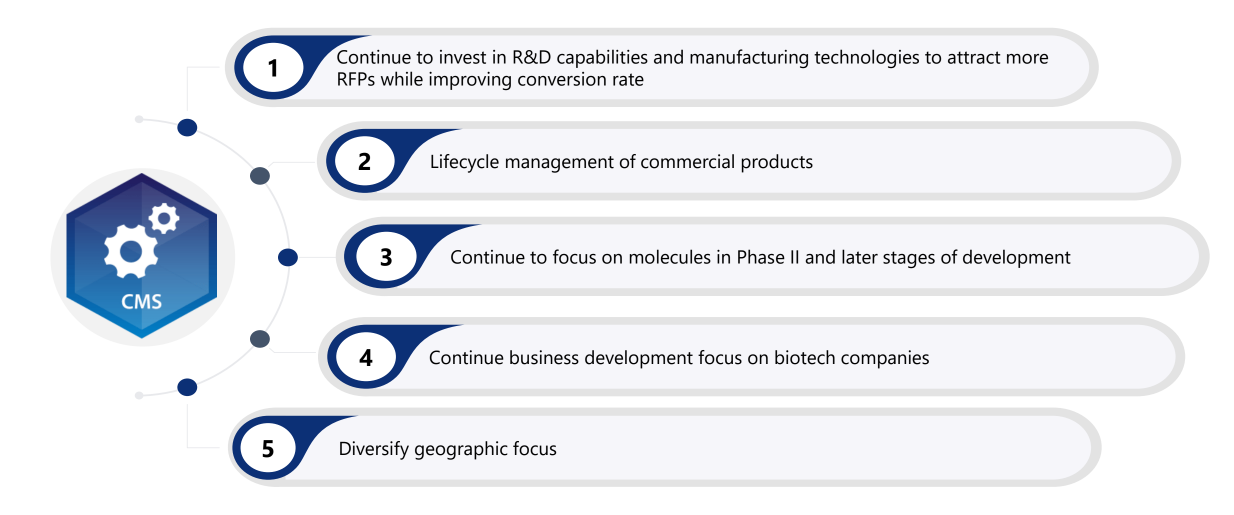
API CDMO Market Size (\$Bn)









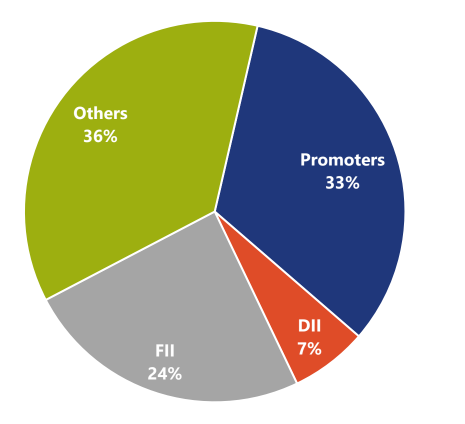




Shareholder Information

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Share Information (as on 31 st Mar 2024)							
NSE Ticker	NEULANDLAB						
BSE Ticker	524558						
Market Cap (Rs. Cr)	8,059						
% free-float	67.20%						
Free-float market cap (Rs. Cr)	5,416						
Shares Outstanding	1,28,29,889						
3M Average Daily Traded Volume (ADTV) (Shares)*	49,699						
3M Average Daily Traded Value (In Rs. Cr)*	30.90						
Industry	Pharmaceuticals						

* Source: BSE & NSE







Particulars (Rs Cr)	Q4FY24	Q4FY23	YoY (%)	Q3FY24	QoQ (%)	FY24	FY23	YoY (%)
Total Income	390.4	415.1	-6.0%	394.9	-1.2%	1,571.1	1,200.9	30.8%
EBITDA	112.2	127.8	-12.2%	122.7	-8.6%	474.5	281.1	68.8%
EBITDA Margin	28.7%	30.8%	-202 bps	31.1%	-233 bps	30.2%	23.4%	680 bps
Profit Before Tax	92.1	110.0	-16.3%	103.8	-11.3%	400.8	215.2	86.2%
PBT Margin	23.6%	26.5%	-290 bps	26.3%	-270 bps	25.5%	17.9%	760 bps
Profit After Tax	67.6	84.5	-20.0%	80.7	-16.2%	299.6	163.1	83.7%
PAT Margin	17.3%	20.4%	-310 bps	20.4%	-310 bps	19.1%	13.6%	550 bps
EPS (Rs.)	52.7	65.9	-20.0%	62.9	-16.2%	233.5	127.1	83.7%

Sustainability at Neuland



Climate change

Reducing greenhouse gas (GHG) emissions intensity and moving towards a balanced portfolio of low carbon energy management

Resource Management

Growing and innovating business solutions through R&D and minimize the use of resources

Local Environmental Protection

Minimizing negative environmental impacts and ensuring the highest standards of EMS

Health & Safety

Making health and safety an integral part of everyday business and culture



People

Creating value and performance culture. Providing work-life balance and engaging employment experience where they can grow and excel

Corporate Governance

Maintain an effective governance and decision-making structure

Ethical Business and Compliance

Fostering an ethical culture and conducting business with integrity and ensure all legal and regulatory compliance

Risk Management

Ensure effective identification of material risks, adequate and effective risk management and internal control

Community

Contributing to the sustainable development of communities through engagement & partnerships and investing in initiatives that make a lasting positive impact

Glossary



Term	Description	Term	Description		
Active Pharmaceutical	Any substance that is intended for incorporation into a finished drug product and is intended to furnish pharmacological activity or other	al activity or other manufacturer			
Ingredient (API)	direct effect in the diagnosis, cure, mitigation, treatment, or prevention of disease, or to affect the structure or any function of the body	Prime APIs	The prime products which typically include mature APIs with relatively higher competition in API space have historically		
Biologic	Biologics can be composed of sugars, proteins, or nucleic acids or complex combinations of these substances, or may be living entities	contributed more than 70% of the total business.			
blologic	such as cells and tissues.	Specialty/ Niche	Molecules in the API space which are complex in nature and are in the nature of 'high value' added products and		
Commercial molecules	Molecules where Neuland is manufacturing for commercial use after the product has been approved	APIs	Neuland's focus has been to develop these molecules from laboratory scale to large commercial quantities		
Custom Manufacturing Solutions (CMS)/ Contract Development and Manufacturing	elopment and Develop and manufacture pharmaceutical ingredients and		Preclinical studies take place in animals before any testing in humans is done.		
Organization (CDMO)		Phase I clinical trial	Researchers test an experimental drug or treatment in a		
Development Molecules	Projects where Phase-3 is over, and molecules have been filed but not yet commercial.	triai	small group of people for the first time.		
DMF	A Drug Master File (DMF) is a submission to the Food and Drug Administration (FDA) that may be used to provide confidential detailed information about facilities, processes, or articles used in the	Phase II clinical trial	The experimental drug or treatment is given to a larger group of people to see if it is effective and to further evaluate its safety.		
	manufacturing, processing, packaging, and storing of one or more human drugs		The experimental study drug or treatment is given to large		
GDS	Generic Drug Substance (GDS) segment which includes Prime products and Specialty products	Phase III clinical trial	groups of people. Researchers confirm its effectiveness, monitor side effects, compare it to commonly used treatments, and collect information that will allow the		
International Council for	Harmonisation project involving regulatory authorities and		experimental drug or treatment to be used safely.		
Harmonisation (ICH) Guidelines	pharmaceutical industry to improve efficiency of new drug development and registration processes		A drug that can enter cells easily because it has a low		
New Chemical Entity (NCE)	NCE is granted to "a drug that contains no active moiety that has been approved by FDA in any other application"	Small molecule products	molecular weight. Once inside the cells, it can affect other molecules, such as proteins, and may cause cancer cells to die. This is different from drugs that have a large molecular weight, which keeps them from getting inside cells easily.		
	Peptides are sequences of molecules called amino acids. Peptides of		Many targeted therapies are small-molecule drugs		
Peptides	precise sequences may occur naturally in the body, but they may also be produced synthetically or using recombinant DNA technology in bacteria and other living systems. These molecules are used to treat a variety of diseases	USFDA	The US Food and Drug Administration is responsible for protecting the public health by ensuring the safety, efficacy, and security of drugs, biological products, and medical devices		

Over 4 decades, Neuland Laboratories Ltd. (BSE:524558, NSE: NEULANDLAB) has been at the forefront of manufacturing APIs through its cGMP manufacturing facilities, working with customers in close to 80 countries. Neuland Labs has developed more than 300 processes and 100 APIs and has filed over 950+ Regulatory filings in the US (67 active US DMFs), the European Union (EU) and other geographies. Its manufacturing facilities are inspected and approved by the U.S. FDA and other leading regulatory agencies. Its record of guality manufacturing and reliability is highlighted by cGMP certifications that include the U.S. FDA, TGA (Australia), EDQM (EU), German Health Authority, ANVISA (Brazil), EMA (EU), Cofepris (Mexico), KFDA (Korea), PMDA (Japan), CFDA (China), FSI "SID &GP" Russia, Health Canada, ISO 9001, ISO14001, OHSAS18001 and ISO 27001.

NEULAND WHERE OPPORTUNITY BECOMES REALITY

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