

March 8, 2023

To	To
Listing Department,	The Corporate Relations Department
NATIONAL STOCK EXCHANGE OF INDIA LIMITED	BSE LIMITED
Exchange Plaza,	Phiroz Jeejeebhoy Towers,
Bandra Kurla Complex, Bandra (E),	25 <sup>th</sup> floor, Dalal Street,
MUMBAI -400 051	<b>MUMBAI -400 001</b>
Company Code No. AUROPHARMA	Company Code No. 524804

Dear Sir/ Madam,

### Sub: Press Release - Eugia Pharma receives USFDA Approval for Lenalidomide Capsules

We enclose a copy of the Press Release that is being issued by the Company in connection with USFDA approval received by Eugia Pharma Specialities Limited, a wholly owned subsidiary of the Company, for Lenalidomide Capsules.

Please take the information on record.

Thanking you,

Yours faithfully, For AUROBINDO PHARMA LIMITED

B. Adi Reddy Company Secretary

Encl: as above

(CIN: L24239TG1986PLC015190)

# AUROBINDO PHARMA LIMITED

www.aurobindo.com

PAN No. AABCA7366H

Corp. Off.: Galaxy, Floors: 22-24, Plot No.1, Survey No.83/1, Hyderabad Knowledge City, Raidurg Panmaktha, Ranga Reddy District, Hyderabad – 500 032, Telangana, India.

Tel: +91 40 6672 5000 / 6672 1200 Fax: +91 40 6707 4044.

Regd. off.: Plot No. 2, Maithrivihar, Ameerpet, Hyderabad-500038 T.S., INDIA Tel: +914023736370/23747340 Fax: +914023741080/23746833 Email: info@aurobindo.com Website: www.aurobindo.com **Press Release** 



Hyderabad, India, March 08, 2023

#### Eugia Pharma receives USFDA Approval for Lenalidomide Capsules

Aurobindo Pharma Limited is pleased to announce that its wholly owned subsidiary company, Eugia Pharma Specialities Limited, has received a final approval from the US Food & Drug Administration (USFDA) to manufacture and market Lenalidomide Capsules, 2.5 mg, 5 mg, 10 mg, 15 mg, 20 mg, and 25 mg, which is bioequivalent and therapeutically equivalent to the Reference Listed Drug (RLD), Revlimid Capsules, 2.5 mg, 5 mg, 10 mg, 15 mg, 20 mg, and 25 mg, of Bristol-Myers Squibb Company. The product is expected to be launched in October 2023 (Volume specific launch).

This is the 155<sup>th</sup> ANDA (including 9 tentative approvals received) out of Eugia Pharma Speciality Group (EPSG) facilities, manufacturing both oral and sterile specialty products.

The approved product is indicated for the treatment of adult patients with Multiple myeloma, in combination with Dexamethasone.

#### About Aurobindo Pharma Limited

Aurobindo Pharma Limited (www.aurobindo.com), (NSE: AUROPHARMA, BSE: 524804, Reuters: ARBN.NS, Bloomberg: ARBP IN) is an integrated global pharmaceutical company headquartered in Hyderabad, India. The Company develops, manufactures, and commercializes a wide range of generic pharmaceuticals, branded specialty pharmaceuticals and active pharmaceutical ingredients globally in over 150 countries.

The company has 24 manufacturing and packaging facilities that are approved by leading regulatory agencies including USFDA, UK MHRA, EDQM, Japan PMDA, WHO, Health Canada, South Africa MCC, Brazil ANVISA. The company's robust product portfolio is spread over 7 major therapeutic/product areas encompassing CNS, Anti-Retroviral, CVS, Antibiotics, Gastroenterological, Anti-Diabetics and Anti-Allergic, supported by a strong R&D set-up.

To know more, please log on to www.aurobindo.com

For further information or queries, please contact:

Soumen Biswas | Deepti Thakur Investor Relations | Corporate Communications Phone: +91 40 66725401 / 66725000 Email: <u>ir@aurobindo.com</u> Disclaimer:

This press release contains statements that may constitute "forward looking statements" including and without limitation, statements relating to product characteristics and uses, sales potential and target dates for product launch, implementation of strategic initiatives, and other statements relating to our future business developments and economic performance. While these forward-looking statements represent our judgment and future expectations concerning the development of our business, a number of risks, uncertainties and other

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factors could cause actual developments and results to differ materially from our expectations. The company undertakes no obligation to publicly revise any forward-looking statements to reflect future events or circumstances and will not be held liable for any use of this information.

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