



GLAND PHARMA LIMITED

July 29, 2026

BSE Limited
Corporate Relationship Department
Phiroze Jeejeebhoy Towers
25th floor, Dalal Street
Mumbai - 400 001
Scrip Code: 543245

National Stock Exchange of India Limited
Listing Department
Exchange Plaza, 5th floor
Plot no. C-1, Block G, Bandra Kurla Complex
Bandra (East), Mumbai - 400 051
Symbol: GLAND (ISIN: INE068V01023)

Dear Sir/Madam,

Sub: Press Release - Gland Pharma receives approval from the United States Food and Drug Administration (USFDA) for Sugammadex Injection 200 mg/2 mL (100 mg/mL) and 500 mg/5 mL (100 mg/mL) Single-Dose Vials

Pursuant to Regulation 30 read with Part A of Schedule III of the Securities and Exchange Board of India (Listing Obligations and Disclosure Requirements) Regulations, 2015; please find attached announcement regarding the receipt of USFDA approval for **Sugammadex Injection 200 mg/2 mL (100 mg/mL) and 500 mg/5 mL (100 mg/mL) Single-Dose Vials**.

This is for your information and records.

Yours truly,

For Gland Pharma Limited

Sampath Kumar Pallerlamudi
Company Secretary & Compliance Officer

Encl: As above

Regd. Office:

Survey No. 143-148, 150 & 151, Near Gandimaisamma 'X' Roads
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Corporate Office:

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Gland Pharma receives approval for Sugammadex Injection 200 mg/2 mL (100 mg/mL) and 500 mg/5 mL (100 mg/mL) Single-Dose Vials.

Hyderabad, July 29, 2026: Gland Pharma Limited (Gland or Company), a generic injectable & ophthalmic-focused pharmaceutical company, has received approval from the United States Food and Drug Administration (USFDA) for its Abbreviated New Drug Application filed for **Sugammadex Injection, 200 mg/2 mL (100 mg/mL) and 500 mg/5 mL (100 mg/mL) Single-Dose Vials.**

The Product is bioequivalent and therapeutically equivalent to the reference listed drug (RLD), BRIDION® Injection of Merck Sharp & Dohme LLC.

The Product is indicated for the reversal of neuromuscular blockade induced by rocuronium bromide and vecuronium bromide in adult and pediatric patients undergoing surgery.

The Company has successfully launched the Product on Day 1 through its marketing partner. According to IQVIA, the product had US sales of approximately USD 1.6 billion for the twelve months ending April 2026.

About Gland Pharma Limited (BSE: 543245, NSE: GLAND)

Gland Pharma was established in 1978 in Hyderabad and has grown over the years from a contract manufacturer of small-volume liquid parenteral products to become one of the largest injectable-focused companies, with a global footprint across 60 countries, including the United States, Europe, Canada, Australia, India, and other markets. It operates primarily under a business-to-business (B2B) model and has an excellent track record in developing, manufacturing, and marketing sterile injectables. It has a wide range of injectables, including vials, ampoules, pre-filled syringes, lyophilized vials, dry powders, infusions, oncology, and ophthalmic solutions. It also enjoys the distinction of having pioneered Heparin technology in India. For more information, log on to: www.glandpharma.com

For investor queries, please mail us at investors@glandpharma.com

This press release may include statements of future expectations and other forward-looking statements based on management's current expectations and beliefs. Actual results may vary materially from those expressed or implied by the statements herein due to economic, business, competitive, technological, and/or regulatory factors changes. Gland Pharma Limited, its directors, and any of the affiliates or employees are under no obligation to, and expressly assume, any obligation to update any particular forward-looking statement contained in this release.