



May 2, 2018

BSE Limited 1st Floor, P J Towers, Dalal Street, Mumbai – 400 001

National Stock Exchange of India Limited Exchange Plaza, 5th Floor, Plot No. C/1, G Block,

Bandra Kurla Complex,
Bandra (East),
Mumbai – 400 051

Re.: Press Release

Dear Sir / Madam,

We enclose herewith a copy of press release dated May 2, 2018 titled "Zydus receives final approval from the USFDA for Bumetanide Tablets USP".

The contents of the press release give full details.

Please bring the aforesaid news to the notice of the members of the exchange and the investors' at large.

Thanking You,

Yours faithfully,

For, CADILA HEALTHCARE LIMITED

UPEN H. SHAH
COMPANY SECRETARY

COM AN DECKERAN

Encl.: As above



Press Release

Press Release

Zydus receives final approval from the USFDA for Bumetanide Tablets USP

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Ahmedabad, 2 May, 2018

Zydus Cadila has received the final approval from the USFDA to market Bumetanide Tablets USP in the strengths of 0.5 mg, 1 mg, and 2 mg. It is used to treat edema associated with congestive heart failure, hepatic and renal disease, including the nephrotic syndrome. It will be manufactured at the group's manufacturing facility at Baddi.

The group now has more than 185 approvals and has so far filed over 320 ANDAs since the commencement of the filing process in FY 2003-04.

About Zydus Cadila

Zydus Cadila is an innovative, global pharmaceutical company that discovers, develops, manufactures and markets a broad range of healthcare therapies. The group employs over 21,000 people worldwide and is dedicated to creating healthier communities globally. Zydus aspires to be a research-based pharmaceutical company by 2020.
