



Dedicated To Life

March 21, 2023

BSE Limited

1st Floor,
P J Towers,
Dalal Street,
Mumbai-400001

Code: 532321

National Stock Exchange of India Limited

Exchange Plaza, 5th Floor,
Plot No. C/1, G Block,
Bandra-Kurla Complex, Bandra (East),
Mumbai-400051

Code: Zyduslife

Re.: Press Release

Dear Sir / Madam,

Please find enclosed a copy of press release dated March 21, 2023 titled **“Zydus receives final and tentative Approval from the USFDA for Tofacitinib Tablets, 5 mg and 10 mg, respectively”**.

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, **ZYDUS LIFESCIENCES LIMITED**

DHAVAL N. SONI
COMPANY SECRETARY

Encl.: As above

Zydus Lifesciences Limited (formerly known as Cadila Healthcare Limited)

Regd. Office : 'Zydus Corporate Park', Scheme No. 63, Survey No. 536, Khoraj (Gandhinagar), Nr. Vaishnodevi Circle,
S. G. Highway, Ahmedabad-382 481, Gujarat, India. | Phone : +91-79-71800000, +91-79-48040000
website : www.zyduslife.com | CIN : L24230GJ1995PLC025878



**Zydus receives final and tentative
Approval from the USFDA for
Tofacitinib Tablets, 5 mg and 10 mg, respectively**

- *Tofacitinib Tablets is also eligible to receive 180 days of shared exclusivity for its 5 mg strength.*

Ahmedabad, India, 21 March, 2023

Zydus Lifesciences Limited's (including its subsidiaries/affiliates, hereinafter referred to as "Zydus") subsidiary Zydus Pharmaceuticals (USA) Inc. has received final approval for Tofacitinib Tablets, 5 mg and tentative approval for Tofacitinib Tablets, 10 mg (USRLD: Xeljanz[®] Tablets) from the United States Food and Drug Administration (USFDA).

Zydus was one of the first ANDA applicants to submit a substantially complete ANDA with a paragraph IV certification for Tofacitinib Tablets, 5 mg and therefore is eligible for 180 days of shared generic drug exclusivity for Tofacitinib Tablets, 5 mg.

Tofacitinib is indicated for the treatment of adult patients with moderately to severely active rheumatoid arthritis and for the treatment of adult patients with active psoriatic arthritis. It is also indicated for the treatment of adult patients with moderately to severely active ulcerative colitis (UC). The drug will be manufactured at the group's formulation manufacturing facility in Moraiya, Ahmedabad (India).

Tofacitinib Tablets, 5 mg and 10 mg had annual sales of USD 900 mn in the United States (IQVIA MAT Dec. 2022).

The group now has 352 approvals and has so far filed over 440* ANDAs since the commencement of the filing process in FY 2003-04.

(*as of 31st December 2022)



**PRESS
RELEASE**

For further information please contact :
The Corporate Communications Department

Zydus Lifesciences Limited
(formerly known as Cadila Healthcare Limited)

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