

JUBILANT PHARMA LIMITED
(Company Registration No. 200506887H)
(Incorporated in the Republic of Singapore)
Registered office: 160 Robinson Road, #17-01 SBF Center, Singapore 068914

August 21, 2017

Singapore Exchange Securities Trading Limited
11 North Buona Vista Drive #06-07
The Metropolis Tower 2
Singapore 138589


Dear Sirs,

We enclose a communication pertaining to USFDA approval.

This is for your kind information and records.

Thanking you,

Yours Faithfully,
For Jubilant Pharma Limited



R. Sankaraiah
Director

Encl.: as above



Jubilant Pharma Ltd.

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Jubilant receives NDA approval for Drax Exametazime

Noida (UP), India, Monday, August 21, 2017

Jubilant Pharma Limited, a material wholly owned subsidiary of Jubilant Life Sciences Ltd, through one of its wholly owned subsidiaries, Jubilant Draximage Inc, Montreal, Canada, has received U.S. Food and Drug Administration (USFDA) approval for **Drax Exametazime** (Kit for the Preparation of Tc99m Exametazime for leukocyte labeling), for its New Drug Application (NDA) pursuant to section 505 (b) (2) filing. This approved new drug application is indicated for leukocyte (white blood cell) labeled scintigraphy as an adjunct in the localization of intra-abdominal infection and inflammatory bowel disease.

This is the sixth approval that we have received from the USFDA during the current financial year. As on June 30, 2017, Jubilant had a total of 84 ANDAs for Oral Solids filed in the US, of which 53 had been approved and 12 Injectable filings of which 9 had been approved.

About Jubilant Pharma Limited

Jubilant Pharma Limited (JPL), a company incorporated under the laws of Singapore and a wholly owned subsidiary of Jubilant Life Sciences Limited, is an integrated global Pharmaceutical company engaged in manufacture and supply of APIs, Solid Dosage Formulations, Radiopharmaceuticals, Allergy Therapy Products and Contract Manufacturing of Sterile and Non Sterile products. JPL has 6 USFDA approved manufacturing facilities in India, US and Canada and R&D centres in India and Canada. The Company has a team of over 3,400 multicultural people across the globe and is committed to deliver value to its customers spread across over 75 countries. It is well recognized as a 'Partner of Choice' by leading pharmaceutical companies globally.

Statements in this document relating to future status, events, or circumstances, including but not limited to statements about plans and objectives, the progress and results of research and development, potential product characteristics and uses, product sales potential and target dates for product launch are forward-looking statements based on estimates and the anticipated effects of future events on current and developing circumstances. Such statements are subject to numerous risks and uncertainties and are not necessarily predictive of future results. Actual results may differ materially from those anticipated in the forward-looking statements. Jubilant may, from time to time, make additional written and oral forward looking statements, including statements contained in the company's filings with the regulatory bodies and its reports to shareholders. The company assumes no obligation to update forward-looking statements to reflect actual results, changed assumptions or other factors.