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# FOSUN PHARMA 复星医药

上海復星醫藥(集團)股份有限公司 Shanghai Fosun Pharmaceutical (Group) Co., Ltd.\*

(a joint stock limited company incorporated in the People's Republic of China with limited liability)

(Stock Code: 02196)

## **OVERSEAS REGULATORY ANNOUNCEMENT**

This announcement is made pursuant to Rule 13.10B of the Rules Governing the Listing of Securities on The Stock Exchange of Hong Kong Limited. The following sets out the "Announcement in Relation to the Approval for Drug Registration of a Subsidiary" published by Shanghai Fosun Pharmaceutical (Group) Co., Ltd.\* (the "Company") on the website of the Shanghai Stock Exchange, for your reference only. The following is a translation of the abovementioned announcement solely for the purpose of providing information. Should there be any discrepancies, the Chinese version will prevail.

By order of the Board

Shanghai Fosun Pharmaceutical (Group) Co., Ltd.\*

Wu Yifang

Chairman

Shanghai, the PRC 13 March 2024

As at the date of this announcement, the executive directors of the Company are Mr. Wu Yifang, Mr. Wang Kexin, Ms. Guan Xiaohui and Mr. Wen Deyong; the non-executive directors of the Company are Mr. Chen Qiyu, Mr. Yao Fang, Mr. Xu Xiaoliang and Mr. Pan Donghui; and the independent non-executive directors of the Company are Ms. Li Ling, Mr. Tang Guliang, Mr. Wang Quandi and Mr. Yu Tze Shan Hailson.

\* for identification purposes only

# Shanghai Fosun Pharmaceutical (Group) Co., Ltd.\* Announcement in Relation to the Approval for Drug Registration of a Subsidiary

The board of directors of the Company and all directors warrant that this announcement does not contain any false information, misleading statement or material omission, and accept legal liability for the truthfulness, accuracy and completeness of the contents herein contained.

#### I. Overview

Shanghai Fosun Pharmaceutical Industrial Development Co., Ltd.\* (上海復星醫藥產業發展有限公司), a subsidiary of Shanghai Fosun Pharmaceutical (Group) Co., Ltd.\* (上海復星醫藥(集團)股份有限公司) (the "Company" and, together with its subsidiaries/units, the "Group", the same applies below), and its shareholding subsidiary Jinzhou Avanc Pharmaceutical Company Limited\* (錦州奧鴻藥業有限責任公司) ("Avanc Pharma"), recently received marketing registration approval from the National Medical Products Administration for the jointly researched and developed Aripiprazole Oral Solution (the "New Drug") for the treatment of schizophrenia in adults and adolescents aged 13-17 years.

# II. General Information of the New Drug

Generic Name: Aripiprazole Oral Solution

Dosage Form: Oral Solution

Specification: 50ml:50mg; 150ml:150mg

Registration Category: Chemicals Category 3

Marketing Authorization Holder/Pharmaceutical Manufacturer: Avanc Pharma

Drug Approval Number: Guo Yao Zhun Zi H20243319, Guo Yao Zhun Zi

# III. Research and Marketing Progress of the New Drug

The New Drug is a chemical drug independently developed by the Group, which is intended to be used primarily for the treatment of schizophrenia.

As of February 2024, the Group has invested approximately RMB 6.12 million (unaudited) in total in the research and development of the New Drug at this stage.

As at the date of this announcement, in addition to the New Drug, other similar products which are already launched in the market in China (excluding Hong Kong, Macau and Taiwan for the purpose of this announcement, the same applies below) mainly include Bosiqing\* (博思清) of Chengdu Kanghong Pharmaceutical Group Co., Ltd.\* (成都康弘藥業集團股份有限公司), Anlvfan(安律凡)® of Zhejiang Otsuka Pharmaceutical Co., Ltd.\* (浙江大塚製藥有限公司) as well as XingPing (醒平)® of Qilu Pharmaceuticals Co., Ltd.\* (齊魯製藥有限公司). According to the latest data of IQVIA CHPA¹, the sales of similar products in China in 2022 was approximately RMB 823 million.

## IV. Impact on the Listed Company and Risk Warning

to varying degrees due to their different sales channels

The approval for marketing registration of the New Drug will bring more options for schizophrenia medication and will further enrich the product line of the Group. It is expected that the approval of the New Drug will not have a material impact on the Group's results at this stage.

Due to the characteristics of the pharmaceutical products industry, the specific sales performance of pharmaceutical products after launch may be affected by factors including, but not limited to, the demand for medication, market competition and sales channels, and is subject to considerable uncertainty. Investors should take note of the investment risks.

<sup>&</sup>lt;sup>1</sup> provided by IQVIA, a provider of professional medical and health information and strategic consultation service in the world; IQVIA CHPA data represents the drug sales market of hospitals with more than 100 beds in China, the actual sales of different drugs may vary from the IQVIA CHPA data

Announcement is hereby made.

Board of Directors of Shanghai Fosun Pharmaceutical (Group) Co., Ltd.\*

13 March 2024

\*For identification purpose only