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山東新華製藥股份有限公司  
**Shandong Xinhua Pharmaceutical Company Limited**

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

(Stock Code: 00719)

**OVERSEAS REGULATORY ANNOUNCEMENT**

This overseas regulatory announcement is made pursuant to Rule 13.10B of the Rules Governing the Listing of Securities on The Stock Exchange of Hong Kong Limited.

Shandong Xinhua Pharmaceutical Company Limited (the “**Company**”) will publish the “Announcement on Having Obtained the Notification of Approval of Supplementary Drug Application and Other Relevant Information” on CNINFO <http://www.cninfo.com.cn> (巨潮資訊網) on 21 January 2025. The English translation of the relevant document is hereby included for reference. If there is any inconsistency between the English version and the Chinese version, the Chinese version shall prevail.

By Order of the Board  
**Shandong Xinhua Pharmaceutical Company Limited**  
**He Tongqing**  
*Chairman*

20 January 2025, Zibo, PRC

As at the date of this announcement, the Board comprises:

Executive Directors:

Mr. He Tongqing (*Chairman*)  
Mr. Xu Wenhui  
Mr. Hou Ning

Non-executive Directors:

Mr. Xu Lie  
Mr. Zhang Chengyong

Independent Non-executive Directors:

Mr. Pan Guangcheng  
Mr. Zhu Jianwei  
Mr. Ling Peixue  
Ms. Cheung Ching Ching, Daisy

## Shandong Xinhua Pharmaceutical Company Limited

### Announcement on Having Obtained the Notification of Approval of Supplementary Drug Application and Other Relevant Information

The Company and its board of directors confirm that the contents of this announcement are true, accurate and complete without any false information, misleading statements or material omissions.

Shandong Xinhua Pharmaceutical Company Limited (hereinafter referred to as “**Xinhua Pharmaceutical**” or the “**Company**”) has recently received the *Notification of Approval of Supplementary Application concerning Drugs* (《药品补充申请批准通知书》) issued by the National Medical Products Administration which approved the supplementary application for the transfer of marketing authorisation holder in relation to the levosalbutamol hydrochloride nebuliser solution (hereinafter referred to as the “**Product**”). Relevant information is now announced as follows:

#### I. Basic information

Drug name:	Levosalbutamol hydrochloride nebuliser solution
Dosage form:	Inhalation preparation
Specification:	Calculated based on C <sub>13</sub> H <sub>21</sub> NO <sub>3</sub> , 3ml: 0.63mg, 3ml: 0.31mg
Drug classification:	Prescription drugs
Applicant:	Shandong Xinhua Pharmaceutical Company Limited
Application matter:	Change of marketing authorisation holder
Reception number:	CYHB2401546、CYHB2401547
Drug approval number:	National Medicine Zhunzi H20243409、 National Medicine Zhunzi H20243408
Notification number:	2024B06214、 2024B06215
Approval Conclusion:	In accordance with the <i>Drug Administration Law of the People’s Republic of China</i> and applicable regulations, upon review, the application concerning the Product complies with applicable requirements for drug registration and the change of the marketing authorisation holder in connection therewith be approved in accordance with the relevant provisions of the <i>Measures for the Administration of Post-marketing Changes of Drugs (Trial)</i> .

#### II. Other relevant information

Xinhua Pharmaceutical and Zhejiang Hengyan Pharmaceutical Technology Co., Ltd. (hereinafter referred to as “**Zhejiang Hengyan**”) entered into a technology transfer contract in December 2022 and January 2024 respectively which stipulates that Zhejiang Hengyan shall make an one-off transfer of its license concerning the marketing and sales of levosalbutamol hydrochloride nebuliser solution and all the rights and interests involved in relevant technology (including production approval documentation, intellectual property rights relating to production technology, commercialisation rights and related rights and benefits etc., including but

not limited to from the aspects of production technology, sales and marketing, etc.) to Xinhua Pharmaceutical. The total technology transfer fee shall be payable by Xinhua Pharmaceutical to Zhejiang Hengyan in accordance with staged instalments as stipulated under the contract. Pursuant to the *Rules Governing the Listing of Shares on Shenzhen Stock Exchange* (《深圳证券交易所股票上市规则》) and the articles of association of the Company (《公司章程》), the present transaction is not required to be submitted for the review and approval of the board of directors or shareholders' meeting of the Company.

The present transaction does not constitute a related party transaction, nor does it constitute a significant asset restructuring as stipulated in the *Measures for Administration of Material Assets Reorganization of Listed Companies* (《上市公司重大资产重组管理办法》).

In December 2024, Xinhua Pharmaceutical submitted supplementary application materials in connection with the change of marketing authorisation holder concerning the Product to the National Medical Products Administration Drug Evaluation Center (CDE) and the application was accepted. In January 2025, Xinhua Pharmaceutical received notification concerning approval of the supplementary application. The conclusion of the review evaluation is that the application for the change of marketing authorisation holder concerning the Product complies with applicable requirements of post-marketing administrative provisions, and the change of marketing authorisation holder concerning the Product was approved.

Levosalbutamol hydrochloride nebuliser solution is used to treat or prevent bronchospasm caused by reversible airway obstructive diseases in adults and adolescents over 6 years old. Levalbuterol is an adrenergic  $\beta$ -receptor agonist, the L-isomer of salbutamol, which can relax bronchial smooth muscle. According to relevant data, sales volume of levosalbutamol inhalation preparations have increased rapidly in recent years. In 2023, sales volume in public hospitals in China amounted to RMB 804 million.

### **III. Impact on the Company and risk warning**

Levosalbutamol hydrochloride nebuliser solution was approved by the National Medical Products Administration in January 2025, and Xinhua Pharmaceutical became the marketing authorisation holder concerning the Product. The launch of this Product is beneficial for enriching our company's respiratory system drug product line, enriching the Company's product line.

The pharmaceutical sales business is susceptible to changes in domestic pharmaceutical industry policies, bidding and procurement processes, changes in the market environment and other factors, and is subject to uncertainty. Investors are advised to invest sensibly and pay attention to investment risks.

By Order of the Board  
**Shandong Xinhua Pharmaceutical Company  
Limited**

20 January 2025