

中华人民共和国 药品GMP证书

CERTIFICATE OF GOOD MANUFACTURING PRACTICES FOR PHARMACEUTICAL PRODUCTS
PEOPLE'S REPUBLIC OF CHINA

证书编号: LN20180012
Certificate No.

企业名称: 沈阳三生制药有限责任公司
Manufacturer: SHENYANG SUNSHINE PHARMACEUTICAL CO., LTD.

地址: 沈阳经济技术开发区十号路1甲3号
Address: No.3 A 1, Road 10, Economy and Technology Development Zone, Shenyang

认证范围: 生物制品[重组人促红素注射液(CHO细胞)(注射剂)、重组人血小板生成素注射液(注射剂)、重组人干扰素 α 2a注射液(小容量注射剂)、注射用重组人干扰素 α 2a(冻干粉针剂)、注射用重组人白介素-2(冻干粉针剂)]
Scope of Inspection: Biological Products [Recombinant Human Erythropoietin Injection (CHO Cell)(Injection), Recombinant Human Thrombopoietin Injection (Injection), Recombinant Human Interferon α 2a Injection (Injection), Recombinant Human Interferon α 2a for Injection (Freeze-dried Powder Injection), Recombinant Human Interleukin-2 for Injection (Freeze-dried Powder Injection)]

经审查,符合中华人民共和国《药品生产质量管理规范》要求。

特发此证。

This is to certify that the above-mentioned manufacturer complies with the requirements of Chinese Good Manufacturing Practices for Pharmaceutical Products.

有效期至 2023 年 5 月 24 日
This certificate remains valid until 24/5/2023

发证机关:
Issued By

辽宁省食品药品监督管理局
Liaoning Food and Drug Administration

Date for Issuing 25/5/2018 2018 年 5 月 25 日

国家食品药品监督管理总局制
CHINA FOOD AND DRUG ADMINISTRATION