



CERTIFICATE OF FDA REGISTRATION

Registration & Listing Of Cosmetic Product Facilities And Products

This certificate only represents that the registered agent has registered the FDA for the following company, and does not represent the FDA endorsement.

Manufacturer Name : **Guangdong Saimei Pharmaceutical Technology Co., Ltd**

Address : First, Second, and Third Floors of Building 5 in Saimei (Guangdong)
Science and Technology Innovation Industrial Park Project, No. 68
Nanxing Road, Shijiao Town, Qingcheng District, Qingyuan,
Guangdong, 511545 China

Type of Business : **MANUFACTURER**

AOKT Technology Service has been registered with the US Food and Drug Administration (FDA) for the aforementioned companies, in compliance with US Food and Drug Administration (FDA). AOKT Technology Service has registered the above facilities with the FDA in accordance with the Modernization of Cosmetics Regulation Act of 2022 (MoCRA) and completed the relevant registration and listing of cosmetic product facilities and products. U.S. FDA does not issue or recognize Certificates of Registration. AOKT Technology Service is not affiliated with the U.S. US Food and Drug Administration (FDA).

Facility FEI Number : **3033358797**

Brand Name : **Jiwancao**

#	Product Listing No.	Product Name	Valid from - until
1	53-038015-573637	Ji Wan Cao Herbal essential oil extracted from plants	11/ 25/ 24 - 11/ 24/ 25

According to the requirements of the Federal Food, Drug, and Cosmetic (FD&C) Act: Manufacturers and processors must register their facilities with the FDA and renew their registration every two years, including any updates to the product and product ingredients. If the FDA determines that cosmetics manufactured or processed by a registered facility have the potential for serious adverse health consequences, the FDA has the authority to suspend the facility's FDA registration.

Note: Under Section 607(c)(5) of the FD&C Act, product updates are required **every year**, including ingredient updates and discontinuation updates. Under Section 607(a)(2) of the FD&C Act, registered facilities must be renewed **every two years**.

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Shenzhen AOKT Technology Service Co., Ltd.

AOKT Technology Service

Web: <http://www.aokt-testing.com/>

Email: aokt-testing@aokt-testing.com

Hery / Registered Agent

Valid from - until
11/ 25/ 2024 - 11/ 24/ 2026