



**Australian Government**

**Department of Health**  
Therapeutic Goods Administration

Dr Xiaolan Kou  
Director- Quality  
Natures Sunshine Products Inc  
1655 North Main Street  
Spanish Fork Utah 84660  
United States of America

Our Reference: 2014/009489

Dear Dr Kou ,

**Subject: Issue of GMP certificate MI-2018-CE-13624-1**

Please find enclosed the GMP certificate for your manufacturing premises.

The certificate remains valid only if re-inspections are conducted when scheduled by the Therapeutic Goods Administration. The inspection frequency is not a reflection of the expiry date shown on the certificate but is consistent with the re-inspection frequency applicable to Australian manufacturers of the same class of products.

The Therapeutic Goods Administration will contact the relevant sponsor/s to arrange the re-inspection of your facility.

Yours sincerely,

Signed and authorised by

Neale Baldwin  
Senior GMP Inspector  
Manufacturing Quality Branch

19 January 2021

Contact: [gmp@tga.gov.au](mailto:gmp@tga.gov.au), phone +61 2 6221 6881 or fax +61 2 6232 8426



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## **Certificate of GMP Compliance of a Manufacturer**

**Certificate Number:**

MI-2018-CE-13624-1

**Issued to:**

Natures Sunshine Products Inc

**Manufacturing Site Address:**

1655 North Main Street  
Spanish Fork Utah 84660  
United States of America

The Therapeutic Goods Administration, the Competent Authority of Australia, confirms that this manufacturer has been inspected following section/s 25(1)(g), 26(1)(g) and/or 26A(3) of the *Therapeutic Goods Act 1989* in connection with marketing authorisation/s listing manufacturers located outside Australia.

From the knowledge gained during inspection of this manufacturer, the latest of which was conducted on 14 to 17 September 2020, it is considered that the manufacturer complies with the Good Manufacturing Practice requirements of the PIC/S Guide to Good Manufacturing Practice for Medicinal Products – 1 July 2018.

This certificate reflects the status of the manufacturing site at the time of the inspection noted above. This certificate remains valid until the expiry date provided that re-inspections are conducted as determined by the Therapeutic Goods Administration as the issuing Authority. This certificate should not be relied upon to reflect the compliance status after the expiry date.

**EXPIRY DATE: 17 December 2022**

**ISSUE DATE: 19 January 2021**

This Certificate remains valid only if re-inspections are conducted when scheduled by the Therapeutic Goods Administration.  
The authenticity of this Certificate may be verified with the Therapeutic Goods Administration as the issuing authority.



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## Certificate of GMP Compliance of a Manufacturer

**Certificate Number:**

MI-2018-CE-13624-1

### MANUFACTURING OPERATIONS

This certificate covers the following steps in the manufacture of therapeutic goods at the manufacturing site address specified above.

Manufacturing Type	Sterility	Dosage Form	Product Category	Manufacturing Step
Medicine manufacture	Non Sterile	Solid Unit Dosage Forms - Tablets	Registered Therapeutic Good	Finished Product Manufacture
Medicine manufacture	Non Sterile	Solid Unit Dosage Forms - Hard Capsules	Listed Therapeutic Good	Finished Product Manufacture
Medicine manufacture	Non Sterile	Liquids	Registered Therapeutic Good	Finished Product Manufacture
Medicine manufacture	Non Sterile	Powder, oral	Listed Therapeutic Good	Finished Product Manufacture
Medicine manufacture	Non Sterile	Herb, dried	Listed Therapeutic Good	Finished Product Manufacture

The following limitations are applicable to these manufacturing operations:

Manufacturing of registered therapeutic goods is limited to complementary medicines only.

This Certificate remains valid only if re-inspections are conducted when scheduled by the Therapeutic Goods Administration.  
The authenticity of this Certificate may be verified with the Therapeutic Goods Administration as the issuing authority.