

# **GMP STATEMENT**

August 30, 2024

### FDA Facility Registration: 14574831328

## OTC Facility FDA Registration: 3004486512

To Whom It May Concern:

We guarantee the products manufactured at **Miramar Cosmetics, Inc DBA MIRAMAR LAB** are in compliance with the Good Manufacturing Practices (GMP) standards. That all ingredient/products delivered are manufactured under Good Manufacturing Practice guidelines following FDA code 21 CFR Part 211.

Products:

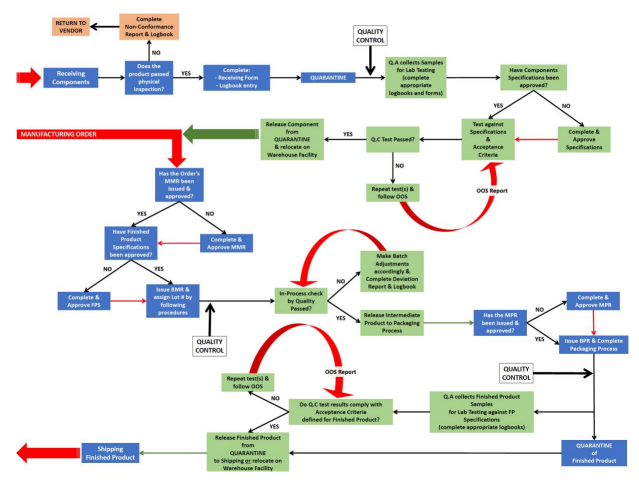
- CISTELA DERMAL OINTMENT 1oz
- CISTELA DERMAL OINTMENT 2oz
- CISTELA DERMAL OINTMENT 4oz
- CISTELA DERMAL OINTMENT 8oz
- CISTELA CREAM 2oz Tube
- CISTELA CREAM 4oz Tube
- CISTELA CREAM 5g SAMPLE TUBE

The above-mentioned facility and controls used for the manufacturing and distribution of **Miramar Cosmetics, Inc DBA MIRAMAR LAB** products are in conformity with Good Manufacturing Practices, in accordance with section 211, title 21, Code of Federal Food, Drugs and Cosmetics Act, and the Amendments thereto.

We hereby certify the above statements to be true and correct to the best of our knowledge and belief.



Manufacturing Process Flow Chart



Sincerely,

Date: 8/30/24

Victor Ramirez CEO Miramar Cosmetics, Inc DBA MIRAMAR LAB

#### 2024

# **CERTIFICATE OF REGISTRATION**

This certifies that:

Miramar Cosmetics Inc DBA Miramar Lab 2301 Nw 107th Ave Ste 101 Doral, FL 33172-2177 United States

is registered with the U.S. Food and Drug Administration pursuant to the Federal Food Drug and Cosmetic Act, as amended by the Bioterrorism Act of 2002 and the FDA Food Safety Modernization Act, such registration having been verified as currently effective on the date hereof by Registrar Corp:

U.S. FDA Registration No.: U.S. FDA UFI (DUNS) No.: U.S. Registration Agent: **14574831328 012873243 Registrar Corp** 144 Research Drive, Hampton, Virginia, 23666, USA Telephone: +1-757-224-0177 • Fax: +1-757-224-0179

This certificate affirms that the above stated facility is registered with the U.S. Food and Drug Administration pursuant to the Federal Food Drug and Cosmetic Act, as amended by the Bioterrorism Act of 2002 and the FDA Food Safety Modernization Act, such registration having been verified as effective by Registrar Corp as of the date hereof, and Registrar Corp will confirm that such registration remains effective upon request and presentation of this certificate until December 31, 2024, unless such registration has been terminated after issuance of this certificate. Registrar Corp makes no other representations or warranties, nor does this certificate make any representations or warranties to any person or entity other than the named certificate holder, for whose sole benefit it is issued. Registrar Corp assumes no liability to any person or entity in connection with the foregoing. The U.S. Food and Drug Administration does not issue a certificate of registration, nor does the U.S. Food and Drug Administration recognize a certificate of registration. Registrar Corp is not affiliated with the U.S. Food and Drug Administration.



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David Lennar

Executive Director Registrar Corp Dated: January 4, 2024 © Copyright 2003-2023 Registrar Corp



# Certificate of Registration 2024

This is to certify that the registration of

MIRAMAR COSMETICS, INC DBA MIRAMAR LAB

2301 NW 107TH AVE, STE 101, DORAL FLORIDA, USA - 33172

with U. S. Food and Drug Administration as required by 21 CFR Part 207 is successfully completed by Miramar Cosmetics, Inc DBA Miramar Lab and verified by Liberty Management Group Ltd.

FEI Number	3004 <mark>486512</mark>
Date of Verification	December 19, 2023
Date of Expiration	December 31, 2024
Certificate Number	2012190123

This certificate does not make representations or warranties to any person or entity other than the named certificate holder; it is issued for record keeping purpose only. This certificate does not denote endorsement or approval of certificate holder's facility or product by the U.S. food and Drug Administration. Liberty management Group Ltd. assumes no liability to any person or entity in connection with the foregoing.

The U.S. Food and Drug Administration does not issue a certificate of registration, nor does the U.S. Food and Drug Administration recognize a certificate of registration. Liberty Management Group Ltd. is not affiliated with the U.S. Food and Drug Administration.



75 Executive Drive, Aurora, Illinois, USA www.fdahelp.us

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Manoj Zacharias President Liberty Management Group LTD. Dated: December 19, 2023 RON DESANTIS, GOVERNOR

MELANIE S. GRIFFIN, SECRETARY

