





Due to a confidentiality agreement signed between Amare Global and the companies that manufacture our products, the names of those companies have been withheld.

UL Verification Services — GMP Inspection Certificate
(UL — Certificate No: i10-34169)

Sunset product is manufactured issued by UL (Underwriters Laboratories), this inspection certificate confirms full compliance with 21 CFR Part 111 — the most current U.S. federal cGMP regulation for dietary supplements. The scope covers manufacturing, packaging, and warehousing of liquid, powder, tablet, capsule, and softgel dosage forms.



UL VERIFICATION SERVICES INC. ISSUES THIS

CERTIFICATE OF INSPECTION

[Blurred Signature]

FOLLOWING INSPECTION OF ITS GOOD MANUFACTURING PRACTICE & QUALITY SYSTEM AND FINDING IT IN CONFORMANCE WITH:

Retail Certification Program Requirements and
ISO 22716:2007 and 2013 FDA Cosmetic Guidelines & ANSI 455-3 GMP for
Cosmetics: 2021

COSMETIC GUIDELINES ON GOOD MANUFACTURING PRACTICES FOR THE FOLLOWING SCOPE OF INSPECTION:

The Manufacturing, Packaging and Warehousing of
Cosmetic Softgel Capsules

Certificate Number: i10-34169-1
Issue Number: 8
Certificate Issue Date: 8/7/2024
Expiration Date: 3/30/2028
Audit Due Date: Decemer 27

Authorized by:

[Signature]

Karen Sak, Business Manager



UL Verification Services Inc.
7036 Snowdrift Road, Suite 200
Allentown, PA 18106
United States of America
800-903-5660



The UL Solutions Logo, Enhanced Certification Mark, and Accreditation Marks indicate satisfactory assessment against the above noted standard / requirements in accordance with the GMP Procedure for Certification, the UL Verification Services LLC Agreement for Use of Certification Symbols, and the scope of assessment. This certificate remains the property of UL Solutions, to whom it must be returned upon request. Revision 4/1/2024



UL VERIFICATION SERVICES INC. ISSUES THIS

CERTIFICATE OF CONFORMANCE

[Blurred Signature]

FOLLOWING ASSESSMENT OF ITS GOOD MANUFACTURING PRACTICE & QUALITY SYSTEM AND FINDING IT IN CONFORMANCE WITH:

Retail Certification Program Requirements and
ISO 22716:2007 and 2013 FDA Cosmetic Guidelines & ANSI 455-3 GMP for
Cosmetics: 2021

COSMETIC GUIDELINES ON GOOD MANUFACTURING PRACTICES FOR THE FOLLOWING SCOPE OF CERTIFICATION:

The Manufacturing, Packaging and Warehousing of
Cosmetic Softgel Capsules

Certificate Number: a10-34169-1
Issue Number: 8
Certificate Issue Date: 8/7/2024
Expiration Date: 3/30/2028
Audit Due Date: Decemer 27

Authorized by:

[Signature]

Karen Sak, Business Manager



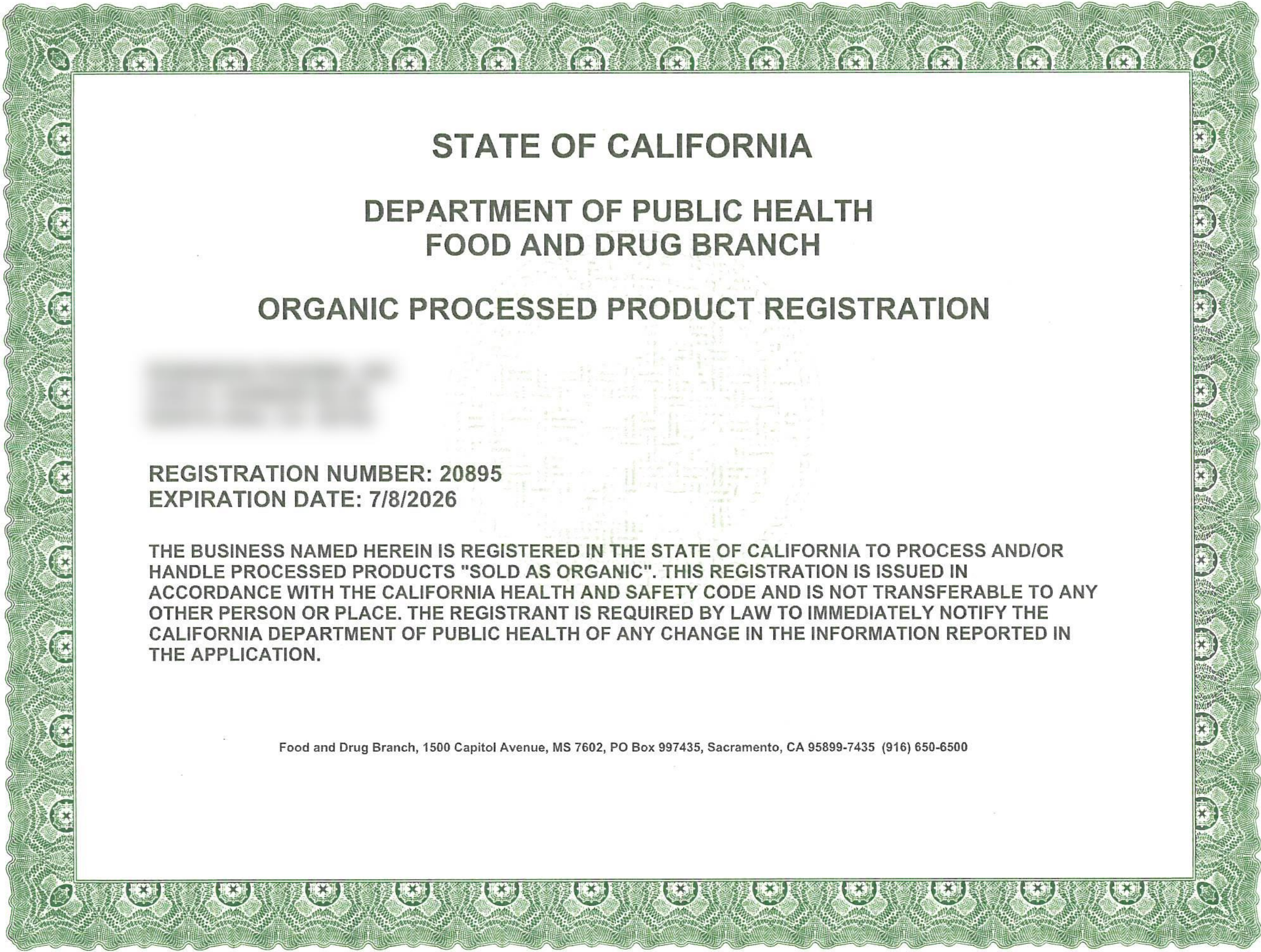
UL Verification Services Inc.
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California Organic Processed Product Registration (California Department of Public Health — No: 20895)

This is from the facility where our **Sunset** product is manufactured registered under the California Organic Food and Farming Act as organically processed products. This registration confirms that those products are legally authorized to carry an organic label under California state law.



The image shows a green-bordered registration certificate from the State of California, Department of Public Health, Food and Drug Branch. The certificate is titled "ORGANIC PROCESSED PRODUCT REGISTRATION" and includes the registration number 20895 and an expiration date of 7/8/2026. A faint background watermark of the state of California is visible. The text states that the business is registered to process and/or handle products "sold as organic" and is not transferable. It also includes the address of the Food and Drug Branch in Sacramento and a disclaimer that the registration is issued in accordance with the California Health and Safety Code.

STATE OF CALIFORNIA
DEPARTMENT OF PUBLIC HEALTH
FOOD AND DRUG BRANCH
ORGANIC PROCESSED PRODUCT REGISTRATION

REGISTRATION NUMBER: 20895
EXPIRATION DATE: 7/8/2026

THE BUSINESS NAMED HEREIN IS REGISTERED IN THE STATE OF CALIFORNIA TO PROCESS AND/OR HANDLE PROCESSED PRODUCTS "SOLD AS ORGANIC". THIS REGISTRATION IS ISSUED IN ACCORDANCE WITH THE CALIFORNIA HEALTH AND SAFETY CODE AND IS NOT TRANSFERABLE TO ANY OTHER PERSON OR PLACE. THE REGISTRANT IS REQUIRED BY LAW TO IMMEDIATELY NOTIFY THE CALIFORNIA DEPARTMENT OF PUBLIC HEALTH OF ANY CHANGE IN THE INFORMATION REPORTED IN THE APPLICATION.

Food and Drug Branch, 1500 Capitol Avenue, MS 7602, PO Box 997435, Sacramento, CA 95899-7435 (916) 650-6500

Printed: 7/11/2025

OSP 07 102308

FDA Food Facility Registration (Dietary Supplement Manufacturer)
(FDA Registration No: 13618051326)

The facility where our **Sunset** product is manufactured, the facility producing our dietary supplement products is registered with the FDA under the Food Safety Modernization Act (FSMA). This registration confirms that the facility's food safety systems meet federal oversight standards.



NSF/GRMA Certificate of Conformance — Dietary Supplement GMP
(NSF International — Certificate No: C0040870)

Issued by NSF International, one of the world's most trusted independent certification bodies, this certificate confirms that our **Sunset** manufacturing facility complies with NSF/ANSI 455-2 — the Good Manufacturing Practices standard for Dietary Supplements.





CERTIFICATE

Audit Number

Issue Date:

Expiration:

First Certification:

Annual Surveillance Date

CERT-000366

AO-000663

03-25-2024

08-06-2026

03-25-2024

01-29-2025

CERTIFICATION

Certification Date:

Expiration:

03-25-2024

08-06-2026



CERTIFICATE OF CONFORMANCE

Facility Type & Description: Campus/Multi-facility, Distributor Onsite, Manufacturing, Primary Facility

Following The Audit Of The Organization's Quality System, Good Manufacturing Practices, And Finding Conformance With:
Global Retailer & Manufacturer Alliance, Inc. Certification Program v1.2
Standard: NSF/ANSI 455-2 - 2021 for Dietary Supplements
Product Types and Manufacturing Stages listed on Page 2
Audit Identification: Onsite



Authorised by David Trosin, Managing Director, Health Sciences Certification

The GRMA Logo, GRMAuditSphere Logo, NSF International's Logo, etc. indicate satisfactory assessment against the Global Retailer & Manufacturer Alliance, Inc.'s Certification Program & NSF/ANSI 455-2 - 2021 for Dietary Supplements. The certification remains the property of NSF International, to whom it must be returned upon request.
Verify Current Certification Status Through Retailer Member Directory

Certificate Issuer

NSF International

789 N. Dixboro Road

Ann Arbor, Michigan, 48105, UNITED STATES

Accreditation # 0216



Page 1 | 3

Valid certificate consists of 3 pages.

Audit Number | AO-000663

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NSF Certificate of Conformity — Dietary Supplement GMP
(NSF/ANSI 455-2:2021)

This is NSF's complementary Conformity Certificate covering an extensive range of product technologies: liquid formula, encapsulation, packaging, labeling, warehousing, mixing, dry formula, coating, and more — all audited and confirmed compliant.



Certificate of Conformity

Print Date

March 02, 2026

Certification Number

C0040870-HSCDS-13

Initial Certification

March 19, 2021

Expiration Date

March 25, 2027

Signed on behalf of
NSF International



David Trosin
Senior Director,
Nutrition and Wellness

NSF International has assessed and confirmed compliance of

Scope: NSF/ANSI 455-2 - 2024
which includes 21CFR Part 111, 21 CFR Part 117, 21 CFR Part 11,
21 CFR Part 1.5 Subpart L & 21 CFR Part 1.9 Subpart O

Product Technologies:
Coating, Dry Formulation, Encapsulation, Liquid Formulation, Mixing,
Packaging/Labeling Operation, Packaging/Labeling Operation - Bulk
Packaging, Packaging/Labeling Operation - Dispensing,
Packaging/Labeling Operation - Primary Packaging,
Packaging/Labeling Operation - Secondary Packaging, Quality Unit
Operations, Warehousing

Product Categories:
Ingestible Liquid, Ingestible Oil, Soft Gel



ANAB
ANSI National Accreditation Board
ACCREDITED
ISO/IEC 17065
PROCESS CERTIFICATION
BODY

NSF International
789 N. Dixboro Road, Ann Arbor, MI 48105 USA

This certificate is the property of NSF International and must be returned upon request.

For the most current and complete information, please access NSF's website (nsf.org)



GMP CERTIFIED
NSF/ANSI 455-2
Dietary Supplements

SQF Food Safety Code — Food Manufacturing Edition 9 (Eurofins Food Assurance — Certificate No: 52985)

This is the facility where our **Sunset** product is manufactured, the SQF (Safe Quality Food) certification is one of the most widely recognized food safety standards in global retail. Product scope: Dietary Supplements (FSC 31) — specifically liquid and powder collagen. Achieved through Unannounced Recertification, meaning production processes are maintained at audit-ready levels at all times.

 Eurofins Food Assurance 2120 Rittenhouse Street, Suite A Des Moines, IA 50321, USA Ph: (515) 299-6979 www.eurofinsus.com/food-safety DATES OF AUDIT: 06/13/2022 - 06/15/2022 NEXT RE-CERTIFICATION DATE: 07/10/2023 DATE OF DECISION: 07/11/2022 EXPIRATION DATE: 09/23/2023 CERTIFICATE NUMBER: 52985 CERTIFICATION TYPE: Unannounced Recertification	Certificate of Registration This acknowledges that is registered as meeting the requirements for the SQF Food Safety Code for Food Manufacturing, Edition 9 Registration schedule Scope of registration [food sector categories and products]: Food sector category: FSC 31: Dietary Supplements Products: Liquid collagen & Powder collagen  Signature of issuing officer Brian Neal Technical Manager  Signature of authorized officer Chuck Russo Vice President  One world. One standard. www.sqf-institute.com SQF Institute is a division of FAMI.  IAS-ANZ G OPS-FM-1055
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Production/quality management certifications of the facility
where our Nitro product is manufactured.



SQF Level 3 Certification

This certification represents the highest distinction attainable from SQF. The SQF Food Safety and Quality Program, formerly known as SQF Level 3 certification, includes all the criteria for levels 1 and 2 SQF certification, while adding defined parameters for achieving consistent quality food grade products.



Current Good Manufacturing Practice Regulations

CGMP refers to Current Good Manufacturing Practice regulations enforced by the FDA. Our CGMP designation assures you of our proper design, monitoring, and control of manufacturing processes and facilities.



NSF

GMP Regulatory Compliance – Good Manufacturing Practices (GMPs) and NSF/ANSI 455 provide a system of processes, procedures and documentation for the dietary supplement, personal care and OTC industries.

Production/quality management certifications of the facility
where our Nitro product is manufactured.



USDA Organic Certification

Our recognized organic certification is your verifiable assurance that we've used only produce grown on soil with no prohibited substances applied for three years before harvesting, as required by USDA organic regulations.



Non GMO

An important distinction and recognized industry standard, non GMO refers to a product produced without genetic engineering or ingredients derived from genetically modified organisms.



Non-GMO Project

Non-GMO Project Verified means that a product is compliant with the Non-GMO Project Standard, which includes stringent provisions for testing, traceability, and segregation. Non-GMO Project Standard is a consensus-based set of criteria crafted with the insight from dozens of industry experts, reflecting a dynamic range of perspectives.

Production/quality management certifications of the facility
where our Nitro product is manufactured.



Gluten Free

Gluten free has become a popular distinction for discerning consumers hoping to achieve improved health, weight loss, and increased energy. Our gluten free distinction lets the consumer know that your products are capable of meeting these strict demands.



Marine Stewardship Council

MSC meets international best practice guidelines and industry standards to ensure that you can trust seafood with the blue MSC label.



Kosher Certification

Kosher certification indicates that a product has been verified by a rabbinic agency to ensure all ingredients, derivatives, tools, and machinery have no trace of non-kosher substances.

Production/quality management certifications of the facility
where our Nitro product is manufactured.



OU Kosher

The OU (Orthodox Union) Kosher is the world's largest and most widely recognized international kosher certification agency. They certify over 1,000,000 products and hold the highest standard of kosher certification. The OU symbol is one of the world's best-known trademarks and offers high quality food standard assurance to customers.



Halal

The word, “halal” means “permissible” in Arabic. This is not the same as kosher. A halal certificate, issued by IFANCA (Islamic Food and Nutrition Council of America), guarantees that a food grade product meets the requirements of Islamic law and has been certified suitable for consumption. Unlike halal, kosher certified foods conform only to Jewish law, and not Islamic.

FDA Drug Establishment Registration / FEI (iLABS Korea Co., Ltd.) (FEI: 3020318393)

The facility where our **Sunrise, Energy+, Edge+, HL5, On Shots, and MentaBiotics** products are manufactured is also registered with the U.S. FDA as a drug manufacturing establishment. This indicates that the quality systems applied at the facility for supplement and cosmetic production are aligned with international pharmaceutical standards.

The certificate is a formal document with a red border and a central seal featuring an eagle. It certifies the registration of iLABS KOREA Co., Ltd. as a cosmetic product facility. The document includes the company's address in Incheon, South Korea, and its facility registration number, 3029936534. It also contains a detailed disclaimer regarding the scope and limitations of the registration and the role of Registrar Corp. as a private registration agent.

2024

CERTIFICATE OF COSMETIC FACILITY REGISTRATION

This certifies that:

iLABS KOREA Co., Ltd.
61 Hambangmoe-ro 401beon-gil, Namdong-gu
Incheon, Incheon 21638
South Korea

has submitted its facility registration statement for the cosmetic product facility identified below with the U.S. Food and Drug Administration pursuant to Chapter VI of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 361 et seq.) and such registration has been verified as currently effective on the date hereof by Registrar Corp.

NAME OF FACILITY: iLABS KOREA Co., Ltd.

FACILITY REGISTRATION NUMBER: 3029936534

This certificate affirms that the above company has filed its Facility Registration Statement for the referenced cosmetic product facility with the U.S. Food and Drug Administration pursuant to the Federal Food, Drug, and Cosmetic Act, as amended by the Modernization of Cosmetics Regulation Act of 2022, (21 U.S.C. 361 et seq.), such registration having been verified as effective by Registrar Corp as of the date hereof. Registrar Corp will confirm that such filing remains effective upon request and presentation of this certificate until December 31, 2024, unless such filing is 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. Registration of a cosmetic product facility or assignment of a registration number does not in any way denote approval by the Food and Drug Administration of the company or the facility. Any representation in labeling or advertising that creates an impression of official approval because of such filing, or such number could be considered misleading and misbranding.

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 a private registration agent not affiliated with the U.S. Food and Drug Administration.

Registrar Corp
144 Research Drive, Hampton, Virginia, 23666, USA
Telephone: +1-757-224-0177 • Fax: +1-757-224-0179
info@registrarcorp.com • www.registrarcorp.com

David Leonard
David Leonard
Executive Director
Registrar Corp
Dated: February 16, 2024

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FDA Cosmetic Facility Registration (iLABS Korea Co., Ltd.) (Registrar Corp)

The facility where our **Sunrise, Energy+, Edge+, HL5, On Shots, and MentaBiotics** products are manufactured, the cosmetic components of our products is registered with the U.S. Food and Drug Administration (FDA). This registration confirms that our manufacturing site operates to auditable standards under U.S. federal law.

FY2024

CERTIFICATE OF REGISTRATION

This certifies that:

ILABS KOREA Co., Ltd.
61 Hambangmoe-ro 401beon-gil, Namdong-gu
Incheon, Incheon 21638
South Korea

is registered with the U.S. Food and Drug Administration for the statutory filing period applicable to U.S. FY 2024 pursuant to part 207 of Title 21, U.S. Code of Federal Regulations.

DUNS Number: **69-418-5577**
FEI: **3020318393**
U.S. Agent/Registrant Contact: **Registrar Corp**
144 Research Drive, Hampton, Virginia, 23666, USA
Telephone: +1-757-224-0177 • Fax: +1-757-224-0179

Filing was performed during the October 1 - December 31, 2023 statutory period, and renewal is not required until the next statutory period of October 1 - December 31, 2024. Registrar Corp will confirm that such registration remains effective upon request and presentation of this certificate, until the end of the year stated above, unless 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. Registration of a drug establishment or drug wholesaler, or assignment of a registration number, or assignment of a NDC number does not in any way denote approval of the firm or its products by the U.S. Food and Drug Administration. Any representation that creates an impression of official approval because of registration or possession of registration number or NDC number is misleading and constitutes misbranding. 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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Telephone: +1-757-224-0177 • Fax: +1-757-224-0179
info@registrarcorp.com • www.registrarcorp.com

David Lennarz
David Lennarz
Executive Director
Registrar Corp
Dated: August 5, 2024

Korean MFDS CGMP Certificate — 우수화장품 제조 및 품질관리기준 (Ministry of Food and Drug Safety, Korea)

The facility where our **Sunrise, Energy+, Edge+, HL5, On Shots, and MentaBiotics** products are manufactured has been certified by the Korean Ministry of Food and Drug Safety as compliant with the "Superior Cosmetic Manufacturing and Quality Management Standards." This is Korea's official CGMP certification.

제 100 호

우수화장품 제조 및 품질관리기준 적합업소 증명서

1. 업소명	주식회사아이엠스코리아
2. 대표자	류치호, 김원태
3. 소재지	인천광역시 남동구 함박외로401번길 61 1층(일부), 3층(일부), 4층(일부), 5층, 인천광역시 남동구 남동대로215번길 43 4층 1층 공장 일부(복관소)
4. 신청분류	천 공장

위의 화장품 제조업소는 「우수화장품 제조 및 품질관리기준」 제30조에 따라 우수화장품 제조 및 품질관리기준 적합업소로 인정되었음을 증명합니다.

2020년 5월 13일

식품의약품안전처장



ISO 14001:2015 — Environmental Management System (Kiwa)

The facility where our **Sunrise, Energy+, Edge+, HL5, On Shots, and MentaBiotics** products are manufactured, holds ISO 14001 certification, an internationally recognized standard for environmental management systems. This demonstrates a systematic approach to waste management, energy consumption, and minimizing environmental impact across production operations.



ISO 9001:2015 — Quality Management System (Kiwa)

The same manufacturer holds ISO 9001 quality management certification. This confirms that all processes — from product development through to manufacturing — are managed in accordance with internationally recognized quality standards.



CERTIFICATE

kiwa

Registration no.	KQ 223155	UNF Code	12
First issue date	2017-08-08	Issue date	2023-08-08
Expiry date	2026-08-08	Renewal date	2023-08-23

Quality Management System Certificate
KS Q ISO 9001:2015/ISO 9001:2015

We certify that the Quality Management System of the Organization:
(주)아이랩스 코리아
is in compliance with the standard KS Q ISO 9001:2015/ISO 9001:2015
for the following products/services:
화장품(스킨케어, 헤어크림, 향수, 바디로션 등, 헤어제품)의 개발 및 제조

General Manager
Seung-Chan Lee


본 인증서에는 KQ 223155 인증번호가 기재되어 있으며, 본 인증서의 유효성을 확인하기 위하여 본 인증서 뒷면에 QR코드가 부착되어 있습니다.

인증 시험장
한국품질인증원 (KQ) 411호
서울특별시 강남구 테헤란로 411호 411호
Tel: +82 2 3387 2101
Fax: +82 2 3387 2106
URL: www.kq.or.kr

한국품질인증원
서울특별시 강남구 테헤란로 411호 411호
Tel: +82 2 3387 2101
Fax: +82 2 3387 2106
URL: www.kq.or.kr

IAF **KAS**

ISO 22716:2007 — Cosmetic GMP (CGMP)
(Kiwa — Certificate KC 22107)

ISO 22716 is the international Good Manufacturing Practice standard for cosmetics. This certificate confirms that good manufacturing practices are fully implemented across skincare, makeup, and hair product production. It is a core requirement for European and global cosmetic markets.



HALAL Certificate

The Halal Certification for Sunrise, HL5, On Shots products is as follows:



KOREA MUSLIM FEDERATION
39 Usadan-ro 10gil, Yongsan-gu, Seoul 04405, Korea
Tel: (82-2) 793-6906, (82-2) 794-7307, Fax: (82-2) 798-9782
www.koreaislam.org

HALAL CERTIFICATE

KMFHC26-280 Date : June 26, 2026

TO WHOM IT MAY CONCERN

Korea Muslim Federation Halal Committee hereby certifies that the listed product below has complied with the requirements from '*OIC/SMIIC 1:2019, MS 1500:2019, OIC/SMIIC 24:2020, OIC/SMIIC 6:2019, OIC/SMIIC 2:2019*' standards which are established according to commonly accepted Sharia(Islamic law).

Scheme of Certification: KMF Halal Food and Beverage Products

Scope of Scheme: Food Products

1. amare SUNRISE(EU)

2. amare SUNRISE(US,SG)

3. amare HL5(EU/TUR)

4. amare ON Shots(EU/TUR)

(Four products only)

Manufacturer : Frombio iksan plant
72, Gukgasikpum-ro, Wanggung-myeon, Iksan-si, Jeonbuk-do, Republic of Korea

Valid until : June 25, 2027



Yours in Islam


Imam Abdul Rahman Lee Ju Hwa
Chairman of Sharia Committee


Ahmad Cho Min Haeng
Chairman of Halal Committee



KOREA MUSLIM FEDERATION
39 Usadan-ro 10gil, Yongsan-gu, Seoul 04405, Korea
Tel: (82-2) 793-6906, (82-2) 794-7307, Fax: (82-2) 798-9782
www.koreaislam.org

HALAL CERTIFICATE

KMFHC26-277 Date : June 12, 2026

TO WHOM IT MAY CONCERN

Korea Muslim Federation Halal Committee hereby certifies that the listed product below has complied with the requirements from '*OIC/SMIIC 1:2019, MS 1500:2019, OIC/SMIIC 24:2020, OIC/SMIIC 6:2019, OIC/SMIIC 2:2019*' standards which are established according to commonly accepted Sharia(Islamic law).

Scheme of Certification: KMF Halal Food and Beverage Products

Scope of Scheme: Food Products

1. Amare Sunrise(TUR,MY)

(One product only)

Manufacturer : Frombio iksan plant
72, Gukgasikpum-ro, Wanggung-myeon, Iksan-si, Jeonbuk-do, Republic of Korea

Valid until : June 11, 2027



Yours in Islam


Imam Abdul Rahman Lee Ju Hwa
Chairman of Sharia Committee


Ahmad Cho Min Haeng
Chairman of Halal Committee

