





**Amare Global ile ürünlerimizi üreten şirketler arasında imzalanan gizlilik anlaşması nedeniyle, bu şirketlerin isimleri açıklanmamıştır.**

# UL Verification Services — GMP Denetim Sertifikası (UL, Sertifika No: i10-34169)

**Sunset** ürünümüzün üretildiği tesise ait, UL (Underwriters Laboratories) tarafından düzenlenen bu denetim sertifikası, 21 CFR Part 111 kapsamında — ABD Takviye Edici Gıda cGMP yönetmeliğinin en güncel versiyonuna — tam uyum sağlandığını belgeler. Sıvı, toz, tablet, kapsül ve softgel formlarındaki ürünlerin üretimi, paketlenmesi ve depolanması denetim kapsamındadır.



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.**  
7036 Snowdrift Road, Suite 200  
Allentown, PA 18106  
United States of America  
800-903-5660

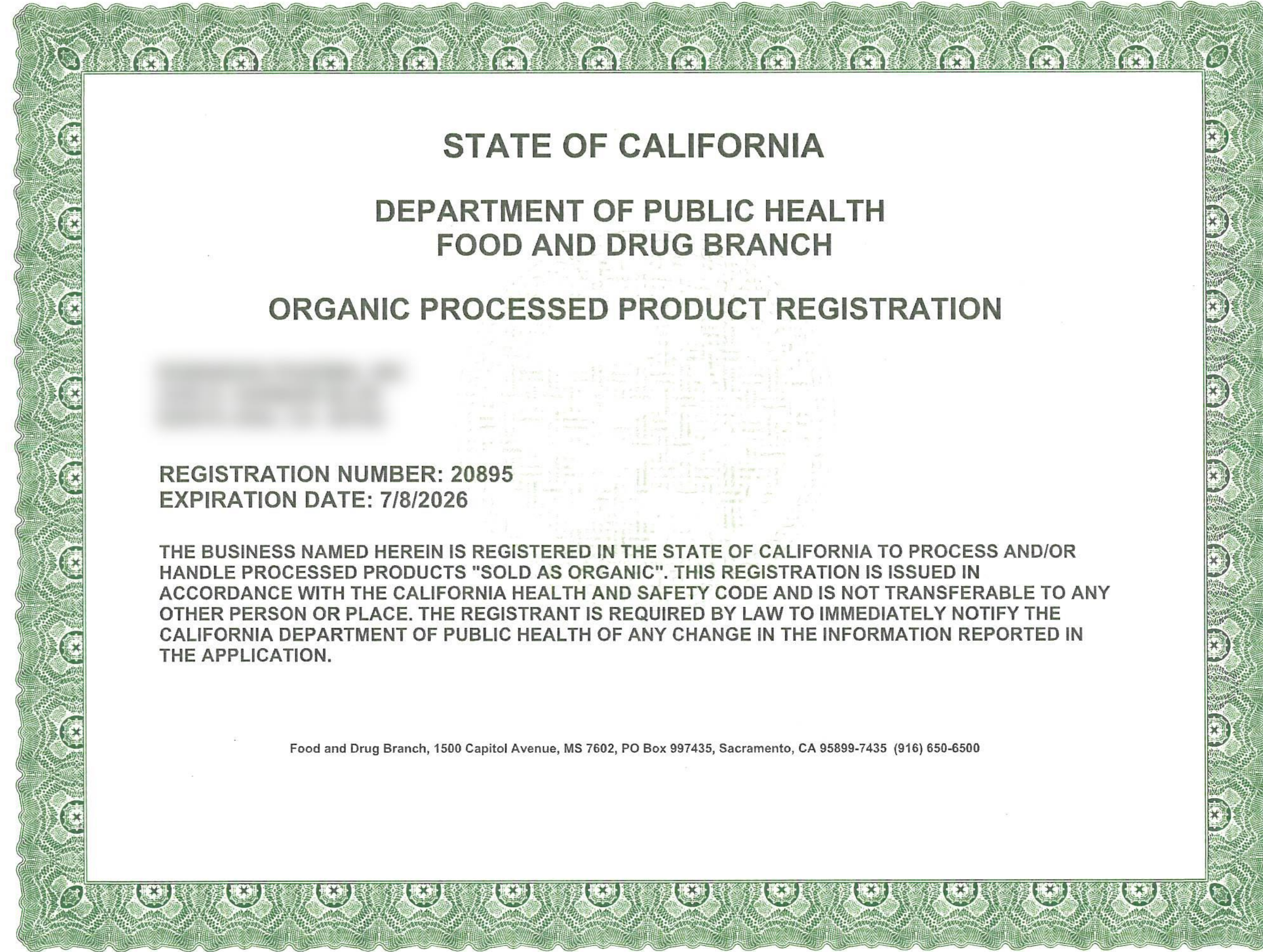


The UL Solutions Logo, Enhanced Certification Mark, and Accreditation Marks indicate satisfactory assessment against the above noted standard / requirements in accordance with the GMP Procedure for Certification, the UL Verification Services LLC Agreement for Use of Certification Symbols, and the scope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

California Organic Processed Product Registration  
(California Sağlık Bakanlığı, No: 20895)

**Sunset** ürünümüzün üretildiği tesise ait olan, California Organik Gıda ve Tarım Yasası kapsamında "organik" olarak işlenmiş ürün kaydına tabidir.

Bu kayıt, söz konusu ürünlerin organik etiketini yasal güvenceyle taşıyabildiğini belgeler.

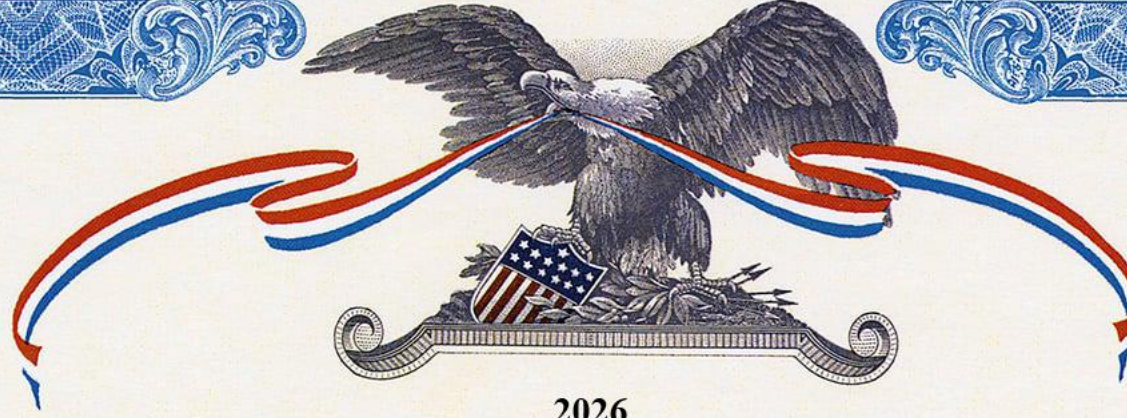


Printed: 7/11/2025

OSP 07 102308

FDA Food Facility Registration (Takviye Gıda Üreticisi)  
(FDA Kayıt No: 13618051326)

**Sunset** ürünümüzün üretildiği tesis, ABD Gıda Güvenliği Modernizasyon Yasası (FSMA) kapsamında FDA'ya kayıtlıdır.  
Bu kayıt, gıda güvenliği sistemlerinin federal denetim standartlarını karşıladığını gösterir.

  
2026

**CERTIFICATE OF REGISTRATION**

*This certifies that:*



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.: **13618051326**  
U.S. FDA UFI (DUNS) No.: **831560578**  
U.S. Registration Agent: **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, 2026, 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.*

  
**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 Lennarz  
Executive Director  
Registrar Corp  
Dated: August 28, 2025  
© Copyright 2003-2023 Registrar Corp

LITHO IN U.S.A.

# NSF/GRMA GMP Uygunluk Belgesi (NSF International, Sertifika No: C0040870)

Bağımsız denetim kuruluđu NSF International tarafından düzenlenen bu belge,  
**Sunset** ürünümüzün üretildiđi tesisimizin NSF/ANSI 455-2 standardına —  
yani Takviye Edici Gıdalar için İyi Üretim Uygulamaları'na — uygun olduđunu onaylar.  
NSF, dünyada en güvenilir üçüncü taraf denetim kuruluşlarından biridir.



|                          |             |
|--------------------------|-------------|
| CERTIFICATE              | CERT-000366 |
| Audit Number             | AO-000663   |
| Issue Date:              | 03-25-2024  |
| Expiration:              | 08-06-2026  |
| First Certification:     | 03-25-2024  |
| Annual Surveillance Date | 01-29-2025  |

|                     |            |
|---------------------|------------|
| CERTIFICATION       |            |
| Certification Date: | 03-25-2024 |
| Expiration:         | 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

  
ANSI National Accreditation Board  
ACCREDITED  
ISO/IEC 17025  
PRODUCT CERTIFICATION  
BODY  
Powered by Intact Platform  
Copyright © 2021 GRMA, All Rights Reserved

## NSF Certificate of Conformity — Takviye Gıda GMP (NSF/ANSI 455-2:2021)

NSF'nin bir önceki belgeyi tamamlayan ikinci Uygunluk Sertifikası'dır. Ürün teknolojileri kapsamı oldukça geniştir: sıvı formül, kapsülleme, ambalajlama, depolama, kaplama ve kuru formül dahil tüm üretim aşamaları denetlenmiştir.



# Certificate of Conformity

Print Date  
March 02, 2026

Certification Number  
C0040870-HSCDS-13

Initial Certification  
March 19, 2021

Expiration Date  
March 25, 2027

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

Signed on behalf of  
NSF International



David Trosin  
Senior Director,  
Nutrition and Wellness



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)



NSF  
GMP CERTIFIED  
NSF/ANSI 455-2  
Dietary Supplements

SQF Food Safety Code — Gıda Üretimi Ed. 9  
(Eurofins Food Assurance, Sertifika No: 52985)

**Sunset** ürünümüzün üretildiği tesise ait SQF (Safe Quality Food) sertifikası, gıda güvenliği alanında küresel perakende sektörünün en yaygın kabul gören belgelerinden biridir. Ürün kategorisi: Takviye Edici Gıdalar (FSC 31) — özellikle sıvı ve toz kolajen. Denetimsiz yeniden sertifikalandırma (Unannounced Recertification) ile elde edilmiştir; bu da üretim süreçlerinin her an denetlenebilir düzeyde olduğu anlamına gelir.



**Eurofins Food Assurance**  
2120 Rittenhouse Street, Suite A  
Des Moines, IA 50321, USA  
Ph: (515) 299-6979  
[www.eurofinsus.com/assurance/food](http://www.eurofinsus.com/assurance/food)

**DATES OF AUDIT:**  
07/17/2025 – 07/18/2025

**NEXT RE-CERTIFICATION DATE:**  
07/10/2026

**DATE OF DECISION:**  
07/29/2025

**EXPIRATION DATE:**  
09/23/2026

**CERTIFICATE NUMBER:**  
FSS202005\_11

**CERTIFICATION TYPE:**  
Announced Recertification

## Certificate of Registration

This acknowledges that

is registered as meeting the requirements for the  
**SQF Food Safety Code for Dietary Supplements Manufacturing, Edition 9**

**Registration schedule**

**Scope of registration [food sector categories and products]:**  
**Food sector category:** FSC 31: Dietary Supplements Manufacturing  
**Products:** Tablets, Capsules, Softgels, Liquids, and Powders

  
**Signature of issuing officer**  
**Brian Neal**

  
One world. One standard.  
SQF Institute is a division of FMI.

  
ANSI National Accreditation Board  
ACCREDITED  
PRODUCT CERTIFICATION BODY  
9166

## Nitro Ürününüzün üretildiği tesise ait üretim/kalite yönetimi sertifikaları



### **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.

## Nitro Ürününüzün üretildiği tesise ait üretim/kalite yönetimi sertifikaları



### 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.

## Nitro Ürününüzün üretildiği tesise ait üretim/kalite yönetimi sertifikaları



### 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.

## Nitro Ürününüzün üretildiği tesise ait üretim/kalite yönetimi sertifikaları



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

**Sunrise, Energy+, Edge+, HL5, On Shots, MentaBiotics** ürünlerimizin kozmetik bileşenlerini üreten tesisimiz,  
ABD Gıda ve İlaç İdaresi'ne (FDA) kayıtlıdır.

Bu kayıt, üretim tesisimizin ABD federal mevzuatı kapsamında denetlenebilir standartlarda faaliyet gösterdiğini belgeler.

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 registration number is 3029936534, and it is valid until December 31, 2024. The document includes a signature of David Leunary, Executive Director of Registrar Corp, dated February 16, 2024. The Registrar Corp's contact information is provided at the bottom left.

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 Leunary  
Executive Director  
Registrar Corp  
Dated: February 16, 2024

©2024-2028 Registrar Corp

FDA Drug Establishment Registration / FEI (iLABS Korea Co., Ltd.)  
(Registrar Corp)

**Sunrise, Energy+, Edge+, HL5, On Shots, MentaBiotics** ürünlerimizin üretildiği tesis, ABD FDA'ya ilaç üretim tesisi olarak da kayıtlıdır. (FEI: 3020318393). Bu, tesisin takviye edici gıda ve kozmetik üretiminde uyguladığı kalite sistemlerini uluslararası ilaç standartlarıyla uyumlu olduğuna işaret eder.

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.

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

Kore MFDS CGMP Sertifikası — 우수화장품 제조 및 품질관리기준  
(Kore Gıda ve İlaç Güvenlik Bakanlığı)

**Sunrise, Energy+, Edge+, HL5, On Shots, MentaBiotics** ürünlerimizin üretildiği tesis,  
Kore Gıda ve İlaç Güvenlik Bakanlığı tarafından "Üstün Kozmetik Üretim ve  
Kalite Yönetimi Standartları"na uygun tesis olarak sertifikalandırılmıştır. Kore'nin resmi CGMP belgesidir.

제 100 호

**우수화장품 제조 및 품질관리기준 적합업소 증명서**

1. 업소명 주식회사아미레스코리아

2. 대표자 유치호, 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 우수  
화장품 제조 및 품질관리기준 적합업소로 인정되었음을 증명합니다.

2020년 5월 13일

식품의약품안전처장



## ISO 14001:2015 — Çevre Yönetim Sistemi (Kiwa)

**Sunrise, Energy+, Edge+, HL5, On Shots, MentaBiotics** ürünlerimizin üretildiği tesis, uluslararası geçerliliğe sahip ISO 14001 standardı kapsamında çevre yönetim sistemi sertifikasına sahiptir.

Bu belge; atık yönetimi, enerji tüketimi ve çevresel etki konularında sistematik bir yaklaşım uygulandığını gösterir.

**CERTIFICATE**

**kiwa**

|                  |            |              |            |
|------------------|------------|--------------|------------|
| Registration no. | KZ 221809  | WF Code      | 12         |
| First issue date | 2017-08-07 | Issue date   | 2023-08-08 |
| Expiry date      | 2026-08-07 | Renewal date | 2023-06-23 |

Environmental Management System Certificate  
**KS I ISO 14001:2015/ISO 14001:2015**

We certify that the Environmental Management System of the Organization:  
**(주)아이랩스 코리아**  
Is in compliance with the standard KS I ISO 14001:2015/ISO 14001:2015  
for the following products/services:  
**화장품(스킨케어, 메이크업, 향수, 바디케어, 헤어케어)의 개발 및 제조**

General Manager  
Seung-Chan Lee

본 인증서 기재 정보에서 인증처의 조직명이나 주소 및 주요적인 사업장임을 증명해 주시기 바랍니다.

**인증 사업장**  
인천광역시 남동구 청학로447번길 41

**키와정보서비스**  
서울특별시 금천구  
영등포로 276  
8층 855호 411호  
Tel: +82-2-2087-2101  
Fax: +82-2-2087-2105  
URL: www.kiwa.kr

**인증 사업장**  
인천광역시 남동구 청학로447번길 41

**IAF** **KAS**  
KAS-00-21

## ISO 9001:2015 — Kalite Yönetim Sistemi (Kiwa)

Aynı üretici, ISO 9001 kalite yönetim sistemi sertifikasına da sahiptir. Kozmetik ürünlerin geliştirilmesinden üretimine kadar tüm süreçlerin uluslararası kalite standartlarına göre yönetildiğini belgeler.



**CERTIFICATE**

**kiwa**

|                  |            |              |            |
|------------------|------------|--------------|------------|
| Registration no. | KQ 225155  | UNF Code     | 12         |
| First issue date | 2017-08-08 | Issue date   | 2020-08-08 |
| Expiry date      | 2026-08-08 | Renewal date | 2025-08-20 |

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 인증서 Kiwa 2000002 인증규정에 따라 발급된 것으로서, 본 인증서의 유효성을 증명하기 위하여 발급되었습니다.

한국표준협회  
서울특별시 강남구  
반포로 279  
6층 615호  
Tel: +82-2-3587-2101  
Fax: +82-2-3587-2106  
URL: www.kfas.or.kr

한국 시험장  
인천광역시 남동구 동막로421번길 51

IAF KAS

## ISO 22716:2007 — Kozmetik GMP (CGMP) (Kiwa)

Uluslararası kozmetik GMP standardı olan ISO 22716 sertifikası, cilt bakımı, makyaj ve saç ürünleri üretiminde iyi imalat uygulamalarının eksiksiz hayata geçirildiğini onaylar. Avrupa ve küresel kozmetik pazarlarında kabul gören temel bir belgedir.



## HALAL SERTİFİKASI

Sunrise, HL5, On Shots Ürünlerimize Ait Halal Sertifikası Aşağıdaki Gibidir:



