

Certificate of Conformance

Certificate
#26-1338822-1

Issue Date
May 27, 2026

Expiration Date
August 27, 2027

Audit Due Date
June 8

Gericare Pharmaceuticals Corporation

1295 Towbin Ave Lakewood, New Jersey 08701 United States

Gericare Pharmaceuticals Corporation

1295 Towbin Ave Lakewood, New Jersey 08701 United States

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

SSCI Technical Benchmarking Requirements (v2020)

Normative Reference Standards Assessed
21 CFR part 111

CURRENT GOOD MANUFACTURING PRACTICE FOR THE FOLLOWING SCOPE OF INSPECTION:

Warehousing, and Distribution of Dietary Supplements

Authorized by:



Karen Sak, Business Manager

UL Verification Services Inc.
2211 Newmarket Pkwy, Suite 106
Marietta, GA 30067
United States of America



Certification Body ID: 21-UL440



The UL Logo and Enhanced Certification Mark indicate satisfactory assessment against the applicable standard in accordance with the SSCI Requirements, the UL LLC Agreement for Use of Certification Symbols, and the scope of assessment. This certificate remains the property of UL, to whom it must be returned upon request. Revision 12/1/2025

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Certificate of Inspection

Certificate

#i23-880382-2

Issue Date

January 6, 2025

Expiration Date

January 23, 2027

Audit Due Date

October 11

Gericare Pharmaceuticals Corporation

1295 Towbin Ave, Lakewood, New York, 08701, United States

Gericare Pharmaceuticals Corporation

1295 Towbin Ave, Lakewood, New York, 08701, United States

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

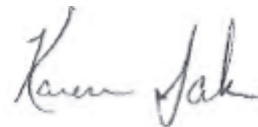
Retail Certification Program Requirements and

Dietary Supplements (21 CFR Part 111 et al., 21 CFR Part 121, 21 CFR Part 117 (Subparts A,B,F as applicable) and 21 CFR Part 1 (Subpart L), Natural Products Association - Good Manufacturing Practices (GMP) Standard for Dietary Supplements)

CURRENT GOOD MANUFACTURING PRACTICE FOR THE FOLLOWING SCOPE OF INSPECTION:

The Packaging and Warehousing of Dietary Supplements in the Form of Tablets, Capsules, Caplets and Gummies

Authorized by:



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 12/11/2024

Certificate of Inspection

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1295 Towbin Ave, Lakewood, New York, 08701, United States	The Packaging and Warehousing of Dietary Supplements in the Form of Tablets, Capsules, Caplets and Gummies
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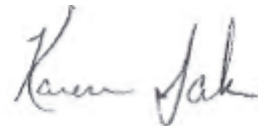
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Authorized by:



Karen Sak, Manager, Client Programs

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7036 Snowdrift Road, Suite 200
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