



**Good
Manufacturing
Practices**

United States Pharmacopeia Quality Systems GMP Audited Certificate

Certificate No.: QSGMP-DSVP-0095 | Original Date: June 5, 2002 | Effective Date: January 1, 2024 | Expiry Date: December 31, 2024

Pharmavite, LLC

1150 Aviation Place
San Fernando, CA 91340

Operates GMP quality systems which meet the requirements set forth in 21 CFR Part 111
*Current Good Manufacturing Practice in Manufacturing, Packaging, Labeling,
or Holding Operations for Dietary Supplements, 21 CFR Part 117 Current Good Manufacturing
Practice, Hazard Analysis and Risk-Based Preventive Controls for Human Food, and USP-NF
general chapter <2750> Manufacturing Practices for Dietary Supplements* for the following
scope:

Dietary Supplement Manufacturing

Holly Chang
Vice President, USP Technical Services

2024

This certificate remains the property of USP and shall be returned immediately upon request. USP issues this GMP certificate for the period stated above or until any major change in the manufacturer's quality systems has taken place, as defined in the license agreement and/or the USP Dietary Supplement Verification Program Manual for Participants. Failure to comply with the provisions of the manual or the license agreement shall render this certificate void, and the right to use the USP Audited Good Manufacturing Practices Mark will be withdrawn. To check the validity of this certificate visit www.uspverified.org.

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Pharmavite, LLC

28310 Livingston Avenue, Valencia, CA 91355
28355 Witherspoon Parkway, Valencia, CA 91355

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