



PERRY JOHNSON LABORATORY ACCREDITATION, INC.

Certificate of Accreditation

Perry Johnson Laboratory Accreditation, Inc. has assessed the Organization of:

Microchem Laboratory, LLC
1700 Chisholm Trail Road, Round Rock, TX 78681

*and hereby declares that the Organization is accredited in accordance with
the recognized International Standard:*

ISO/IEC 17025:2017
**& Meets the Requirements of Good Laboratory Practice for Nonclinical
Laboratory Studies, Title 21
CFR Part 58 Accreditation Program**

Biological Testing
(As detailed in the supplement)

Accreditation claims for conformity assessment activities shall only be made from the addresses referenced within this certificate and shall apply solely to those activities identified in the related scope. This Accreditation is granted subject to the Accreditation Body rules governing the Accreditation referred to above, and the Organization hereby commits to observing and complying with those rules in their entirety.

For PJLA:

Initial Accreditation Date:

Issue Date:

Expiration Date:

February 26, 2025

February 26, 2025

April 20, 2025

Tracy Szerszen
President

Accreditation No.:

Certificate No.:

Extension Date:

128395

L25-160

July 28, 2025

*The validity of this certificate is maintained through ongoing assessments based
on a continuous accreditation cycle. The validity of this certificate should be
confirmed through the PJLA website: www.pjllabs.com*

Perry Johnson Laboratory
Accreditation, Inc. (PJLA)
755 W. Big Beaver, Suite 1325
Troy, Michigan 48084

| FIELD OF TEST | ITEMS, MATERIALS, OR PRODUCTS TESTED | COMPONENT, CHARACTERISTIC, PARAMETER TESTED | SPECIFICATION OR STANDARD METHOD | TECHNOLOGY OR TECHNIQUE USED | FLEX CODE | LOCATION OF ACTIVITY |
|---------------|--------------------------------------|---|----------------------------------|------------------------------|-----------|----------------------|
| Biological | Reusable Medical Devices | Validation of Cleaning Efficacy | ANSI/AAMI ST 98 | Spectrophotometer | F1, F2 | F |
| Biological | Reusable Medical Devices | Validation of Sterilization/Reprocessing | AAMI TIR 12 | Plating/Incubator | F1, F2 | F |
| Biological | Cosmetics | Evaluation of Antimicrobial Protection of Cosmetic Products | ISO 11930 | Plating/Incubator | F1, F2 | F |

1. Location of activity:

Location Code

| | |
|---|--|
| F | Conformity assessment activity is performed at the CABs fixed facility |
| O | Conformity assessment activity is performed onsite at the CABs customer location |
| M | Conformity assessment activity is performed from a mobile facility |

Location

2. Flex Code:

F0- Fixed scope item. No deviations allowed to the line item as identified, except for updating to the most recent version of an accredited standard method after verification.

F1- Laboratory has the capability to test a new item, material, matrix, or product similar in composition to item, material, matrix, or product identified on the scope

F2- Laboratory has the capability to introduce the newest revision of an accredited authoritative standard method (with no modifications) identified on the scope

F3- Laboratory has the capability to introduce a parameter/component/analyte to an accredited test method identified on the scope

F4- Laboratory has the capability to introduce a new revision of an accredited non-standard method using the same technology or technique identified on the scope

F5- Laboratory has the capability to introduce a validated method that is equivalent to an accredited method (using same technology or technique) identified on the scope

3. The above scope of accreditation was created based on a former ILAC MRA Signatory's certificate policy. Based on the intent of the ILAC MRA, PJLA recognizes other scopes issued by other ILAC signatories. This scope will be modified based on PJLA's Policy following the next on-site assessment.