



Accredited Laboratory

A2LA has accredited

HIGHPOWER VALIDATION TESTING & LAB SERVICES INC.

Rochester, NY

for technical competence in the field of

Biological Testing

This laboratory is accredited in accordance with the recognized International Standard ISO/IEC 17025:2017 *General requirements for the competence of testing and calibration laboratories*. This accreditation demonstrates technical competence for a defined scope and the operation of a laboratory quality management system (refer to joint ISO-ILAC-IAF Communiqué dated April 2017).



Presented this 20th day of December 2022.

A blue ink signature of Mr. Trace McInturff, written over a horizontal line.

Mr. Trace McInturff, Vice President, Accreditation Services
For the Accreditation Council
Certificate Number 3718.01
Valid to December 31, 2024

For the tests to which this accreditation applies, please refer to the laboratory's Biological Scope of Accreditation.



SCOPE OF ACCREDITATION TO ISO/IEC 17025:2017

HIGHPOWER VALIDATION TESTING & LAB SERVICES INC.

125 Highpower Road

Rochester, NY 14623

Brandon Taylor Phone: (585) 743-1930

BIOLOGICAL

Valid To: December 31, 2024

Certificate Number: 3718.01

In recognition of the successful completion of the A2LA evaluation process, accreditation is granted to this laboratory to perform the following tests on reusable medical devices, medical devices, biological indicators, chemical indicators, and medical device raw materials:

<u>Test</u>	<u>Test Method(s)</u>
Biological Indicators – D-Value Verification	USP <55>; ANSI/AAMI/ISO 11138-1
Chemical Indicators – Compliance Testing (Dry Heat, Steam)	ANSI/AAMI/ISO 11140-1
Materials Compatibility	AAMI TIR 12; AAMI TIR 17
Sterilization Packaging Systems – Accelerated Aging of Sterile Barrier Systems for Medical Devices	ASTM F1980
Sterilization Packaging Systems – Detecting Seal Leaks in Porous Medical Packaging by Dye Penetration	ANSI/AAMI/ISO 11607-1; AAMI/ISO TIR 16775; ASTM F1929
Sterilization Packaging Systems – Dry Time Testing	ANSI/AAMI ST77; AAMI TIR 12
Sterilization Packaging Systems – Real Time Shelf Life	ANSI/AAMI/ISO 11607-1; AAMI/ISO TIR 16775
Sterilization Packaging Systems – Sterilant Penetration / Thermal Profile	ANSI/AAMI ST77; AAMI TIR 12
Sterilization Packaging Systems – Whole Package Integrity Test	ANSI/AAMI/ISO 11607-1; AAMI/ISO TIR 16775; ANSI/AAMI ST77
Validation of Cleaning Efficacy of Reusable Medical Devices	AAMI ST 98
Validation of Sterilization	ANSI/AAMI/ISO 14937; ANSI/AAMI/ISO 17665-1; AAMI TIR 12
Validation of Thermal Disinfection of Reusable Medical Devices	ISO 15883-1, ISO 15883-2



*Joint ISO-ILAC-IAF
Communique on the
Management Systems Requirements of ISO/IEC 17025,
General Requirements for the competence of testing and
calibration laboratories*

*A laboratory's fulfillment of the requirements of ISO/IEC 17025 means the laboratory meets both the technical competence requirements and **management system requirements** that are necessary for it to consistently deliver technically valid test results and calibrations. The **management system requirements** in ISO/IEC 17025 are written in language relevant to laboratory operations and operate generally in accordance with the principles of ISO 9001.*

A handwritten signature in blue ink, appearing to read "L. Ndri", written over a horizontal line.

ISO Acting Secretary General

A handwritten signature in black ink, reading "Mats Malmqvist Nilsson", written over a horizontal line.

ILAC Chair

A handwritten signature in black ink, consisting of stylized Chinese characters, written over a horizontal line.

IAF Chair