

# **Certification and Documentation Packet**

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#### SCOPE OF ACCREDITATION TO ISO/IEC 17025:2017

### NELSON LABORATORIES BOZEMAN, LLC1 1765 South 19th Avenue Bozeman, MT 59718

Dr. Margret Butler Phone: 406-587-5735 ext. 163 Danielle Goveia Phone: 406-587-5735 ext. 136

### **BIOLOGICAL**

Valid To: May 31, 2024 Certificate Number: 3945.01

In recognition of the successful completion of the A2LA evaluation process, (including an assessment of the laboratory's compliance to U.S. Food and Drug Administration's GLP (Good Laboratory Practices Act) requirements as specified in the Code of Federal Regulations Title 21 part 58 (21CFR58)), accreditation is granted to this laboratory to perform In Vivo and In Vitro testing for qualitative and quantitative analysis of medical devices, textiles, wound dressings, respirators, gloves, condoms, and sanitizing substances, including the antimicrobial properties of soaps, hand sanitizers, preoperative skin preparations, and disinfectants.

Test Technology	Test Method		
Clinical Bactericidal Tests - In Vivo			
Chemical Disinfectants and Antiseptics - Hygienic Hand Rub - Test Method and Requirements (Phase 2/Step 2)	EN1500		
Chemical Disinfectants and Antiseptics - Hygienic Handwash - Test Method and Requirements (Phase 2/Step 2)	EN 1499		
Standard Test Method for Determining the Bacteria - Eliminating Effectiveness of Hygienic Handwash and Hand Rub Agents Using the Finger Pads of Adults	ASTM E2276		
Standard Test Method for Determining the Bacteria - Eliminating Effectiveness of Healthcare Personnel Hand Rub Formulations Using Hands of Adults	ASTM E2755		
Standard Test Method for Evaluation of Pre-operative, Pre-catheterization, or Pre-injection Skin Preparations	ASTM E1173		
Standard Test Method for Evaluation of Surgical Hand Scrub Formulations	ASTM E1115		
Standard Test Method for Evaluation of the Effectiveness of Handwash Formulations Using the Paper Towel (Palmar) Method of Hand Contamination	ASTM E2784		

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Test Technology	Test Method
Standard Test Method for Evaluation of the Effectiveness of Health Care Personnel Handwash Formulations	ASTM E1174
Standard Guide for Evaluation of Residual Effectiveness of Antibacterial Personal Cleansing Products	ASTM E2752
Clinical Virucidal Tests - In Vivo	
Standard Test Method for Determining the Virus - Eliminating Effectiveness of Hygienic Handwash and Hand Rub Gents Using the FingerPads of Adults	ASTM E1838
Standard Test Method for Evaluation of Hygienic Handwash and Hand Rub Formulations for Virus - Eliminating Activity Using the Entire Hand	ASTM E2011
Clinical Respirator Fit Test - In Vivo	
Standard Test Method for Respirator Fit Capability for Negative-Pressure Half-Facepiece Particulate Respirators	ASTM F3407
Bactericidal Tests and Fungicidal - In Vitro	
Chemical Disinfectants and Antiseptics. Quantitative Suspension Test for the Evaluation of Basic Bactericidal Activity of Chemical Disinfectants and Antiseptics. Test Method and Requirements (Phase 1)	EN 1040
Chemical Disinfectants and Antiseptics. Quantitative Suspension Test for the Evaluation of Bactericidal Activity of Chemical Disinfectants and Antiseptics Used in Food, Industrial, Domestic and Institutional Areas. Test Method and Requirements (Phase 2, Step 1)	EN 1276
Chemical Disinfectants and Antiseptics - Quantitative Suspension Test for the Evaluation of Bactericidal Activity in the Medical Area - Test Method and Requirements (Phase 2, Step 1)	EN 13727
Chemical Disinfectants and Antiseptics - Quantitative Suspension Test for the Evaluation of Mycobactericidal Activity of Chemical Disinfectants in the Medical Area including Instrument Disinfectants - Test Method and Requirements (Phase 2, Step 1)	EN 14348
Chemical Disinfectants and Antiseptics. Quantitative Suspension Test for the Evaluation of Fungicidal or Yeasticidal Activity in the Medical Area. Test Method and Requirements (Phase 2, Step 1)	EN 13624



Test Technology	Test Method
Chemical Disinfectants and Antiseptics. Quantitative Suspension Test for the Evaluation of Basic Fungicidal or Basic Yeasticidal Activity of Disinfectants and Antiseptics. Test Method and Requirements (Phase 1)	EN 1275
Chemical Disinfectants and Antiseptics - Quantitative Suspension Test for the Evaluation of Fungicidal or Yeasticidal Activity of Chemical Disinfectants and Antiseptics used in Food, Industrial, Domestic and Institutional Areas - Test Method and Requirements (Phase 2, Step 1)	EN 1650
Chemical Disinfectants and Antiseptics. Quantitative Test for the Evaluation of Bactericidal and Yeasticidal and/or Fungicidal Activity of Chemical Disinfectants in the Medical Area on Non-Porous Surfaces without Mechanical Action. Test method and requirements (Phase 2, Step 2)	EN 17387
Chemical Disinfectants and Antiseptics-Quantitative Test Method for the Evaluation of Bactericidal and Yeasticidal Activity on Non-porous Surfaces with Mechanical Action Employing Wipes in the Medical Area (4 - field test) - Test Method and Requirements (Phase 2, Step 2)	EN 16615
Chemical Disinfectants and Antiseptics – Quantitative Non-porous Surface Test for the Evaluation of Bactericidal and/or Fungicidal Activity of Chemical Disinfectants Used in Food, Industrial, Domestic, and Institutional Areas-Test Method, and Requirements Without Mechanical Action (Phase 2, Step 2)	EN 13697
Measurement of Antibacterial Activity on Plastics and Other Non-porous Surfaces	ISO 22196
Standard Test Method for Assessment of Antimicrobial Activity for Water Miscible Compounds Using a Time - Kill Procedure	ASTM E2783
Germicidal and Detergent Sanitizing Action of Disinfectants	AOAC 960.09
Antimicrobial Effectiveness Testing	USP <51>
Virucidal Tests - <i>In Vitro</i>	
Chemical Disinfectants and Antiseptics - Quantitative Suspension Test for the Evaluation of Virucidal Activity in the Medical Area - Test Method and Requirements (Phase 2, Step 1) (Includes Amendment:2019)	EN 14476
Chemical Disinfectants and Antiseptics. Quantitative Non-Porous Surface Test without Mechanical Action for the Evaluation of Virucidal Activity of Chemical Disinfectants used in the Medical Area. Test Method and Requirements (Phase 2, Step 2)	EN 16777



Test Technology	Test Method						
Measurement of Antiviral Activity on Plastics and Other Non-porous Surfaces	ISO 21702						
Textiles - Determination of Antiviral Activity of Textile Products	ISO 18184						
Standard Practice to Assess the Activity of Microbicides Against Viruses in Suspension	ASTM E1052						
Standard Practice to Assess Virucidal Activity of Chemicals Intended for Disinfection of Inanimate, Non-porous Environmental Surface	ASTM E1053						
Neutralization of Antimicrobial Activity - In Vitro							
Standard Practices for Evaluation of Inactivators of Antimicrobial Agents	ASTM E1054						
Wound Dressing Evaluation – In Vitro							
Test Method for Primary Wound Dressings – Part 1: Aspects of Absorbency	EN 13726-1 Section 3.2 & 3.3						
Test Methods for Primary Wound Dressings – Part 2: Moisture Vapor Transmission Rate of Permeable Film Dressings	EN 13726-2						

<sup>&</sup>lt;sup>1</sup>This scope meets the A2LA P112 Flexible Scope Policy.



# **Accredited Laboratory**

A2LA has accredited

# NELSON LABORATORIES BOZEMAN, LLC

Bozeman, MT

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 laboratory also meets the U.S. Food and Drug Administration's GLP (Good Laboratory Practices Act) requirements as specified in the Code of Federal Regulations Title 21 part 58 (21CFR58). 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 19th day of July 2022.

Vice President, Accreditation Services For the Accreditation Council Certificate Number 3945.01 Valid to May 31, 2024

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

Registrant - Nelson Laboratories Bozeman, LLC (787813161)

Contact Address: 1765 S. 19th Avenue

Danielle Goveia City, State, Zip:Bozeman, MT, 59718
Country: USA

Telephone Number Email Address

City, State, Zip:Bozeman, MT, 59718
Country: USA

dgoveia@nelsonlabs.com

Facility

Name	Address	ID/FEI	Business Operations
Nelson Laboratories Bozeman, LLC	Address: 1765 S. 19th Avenue City, State, Zip:Bozeman, MT, 59718 Country: USA	787813161/1000221600	clinical bioequivalence or bioavailability study, in vitro bioequivalence or bioanalytical testing
Contact	Address	Telephone Number	Email Address
Danielle Goveia	Address: 1765 S. 19th Avenue City, State, Zip:Bozeman, MT, 59718 Country: USA	+406-587-5735;ext=136	dgovcia@nelsonlabs.com

Revised: 12/2023

Registrant - Nelson Laboratories Bozeman, LLC (787813161) Contact

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dgoveia@nelsonlabs.com

analysis

Establishment

Nelson Laboratories Bozeman, LLC

Danielle Goveia

ID/FEI Address **Business Operations** Name Address: PO Box 190

City, State, Zip:Bozeman, MT, 59718

Country:

787813161/1000221600

Address Telephone Number Email Address

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Revised: 12/2023

Contact

Danielle Goveia

### DEPARTMENT OF HEALTH AND HUMAN SERVICES PUBLIC HEALTH SERVICE

#### FOOD AND DRUG ADMINISTRATION

ESTABLISHMENT REGISTRATION AND LISTING FOR HUMAN CELLS, TISSUES, AND CELLULAR AND TISSUE-BASED PRODUCTS DESCRIBED IN 21 CFR 1271.10

Ext.:

FEI: 3026934915

Other FDA Registrations:

Blood: Devices:

Drugs: FEI: 1000221600

Reason For Last Submission: Annual Registration/Listing

Last Annual Registration Year: 2024
Last Registration Receipt Date: 12/22/2023
Summary Report Print Date: 01/05/2024

Legal Name and Location:

Bozeman, Montana 59718

Phone: 406-587-5735

Nelson Laboratories Bozeman, LLC

1765 S. 19th Ave

USA

Robert Thoreson, Director of Quality Assurance 6280 South Redwood Road

Salt Lake City, Utah 84123

Reporting Official:

USA

Phone: 801-290-7618 Ext. rthoreson@nelsonlabs.com

Satellite Recovery Establishment:

Parent Manufacturing Establishment FEI No.:

Testing For Micro-Organisms Only: Yes

Note: FDA acceptance of an establishment registration and HCT/P listing does not constitute a determination that an establishment is in compliance with applicable rules and regulations or that the HCT/P is licensed or approved by FDA (21 CFR 1271.27(b)).

No

	Establishment Functions											
HCT/P(s) Donor Type	Donor Type(s)	Recover	Screen	Donor Testing	Package	Process	Store	Label	Distribute	Date of Discontinuance	Date of Resumption	Proprietary Name(s)
Amniotic Membrane						Х						
Blood Vessel						Х						
Bone						Х						
Cardiac Tissue - non-valved						Х						
Cartilage						Х						
Cornea						Х						
Dura Mater						Х						
Embryo												
Fascia						Х						
Heart Valve						Х						
HPC Apheresis												
HPC Cord Blood												
Ligament						Х						
Nerve Tissue						Х						
Oocyte												
Ovarian Tissue						Х						
Pancreatic Islet Cells - autologous						Х						
Parathyroid						Х						
Pericardium						Х						
Peripheral Blood Mononuclear Cells												
Peritoneal Membrane						Х						
Sclera						Х						
Semen												
Skin						Х						
Tendon						Х						
Testicular Tissue						Х						
Tooth Pulp						Х						
Umbilical Cord Tissue						Х						

Additional	Information:	No addit	ic

No additional information provided.

Proprietary Name(s):

FEI: 3026934915

Legal Name:

Nelson Laboratories Bozeman, LLC

New Search Results Return to: Search Results

## IRB Organization Information

IORG0004971 - Nelson Laboratories Bozeman, LLC (Active)

Located at: Bozeman, MONTANA

**Expires**: 05/05/2025

### IRBs for this Organization: 1

**Agency Only Access** 

IRB# IRB Name City State/Country Status IRB Type
IRB00005939 Gallatin IRB #1 Bozeman MONTANA Active OHRP/FDA

Department of Health and Human Services (DHHS) | Office for Human Research Protections (OHRP) | Accessibility | HHS Vulnerability Disclosure