

# BIODEX

## DECLARATION OF CONFORMITY

### PRODUCT IDENTIFICATION

Type of Equipment	Product Name(s)	Model/Number(s)
Radionuclide Dose Calibrators	Atomlab 500	086-330/-331/-332/-336

Serial Number N/A

### MANUFACTURER

Name of Company	Address	Representative
Biodex Medical Systems, Inc.	20 Ramsey Road Shirley, NY 11967-4704 USA	Clyde Schlein <i>Vice President, Regulatory Affairs &amp; Compliance</i>

### AUTHORIZED REPRESENTATIVE

Name of Company	Address	Telephone/Email
Emergo Europe	Prinsessegracht 20 2514 AP The Hague The Netherlands	Phone: +31.70.345.8570 Fax: +31.70.346.8570

### REGISTRATION INFORMATION

Notified Body and ID#	CE Certificate No & Expiration Date	Date CE Marking First Applied
Intertek Services	41312458-01 7-20-2023	N/A

### CONFORMITY ASSESSMENT

Device Classification	Route to Compliance	Standards Application
Class I Measuring Rule #12 Annex V	Annex V of MDD 93/42/EEC Council Directive	EN60601-1 Ed3 EN 60601-1-2:2007 IEC 60601-1-1:2000 Ed 2 IEC 601-1:1988 +A1 +A2 UL 60601-1:2003 CAN/CSA c22.2 No.:601-1-M90 IEC 60601-1-4:2000

### QUALITY SYSTEM REGISTRAR

Intertek Services Certified Quality Management System:  
ISO 13485:2016 Certificate: 0084059  
Biodex Medical Systems, Inc. declares the above mentioned products meet the provision of the Council Directive 93/42/EEC for Medical Devices.

**Company Representative:** Clyde Schlein  
**Title:** Vice President, Regulatory Affairs & Compliance  
**Date:** May 1, 2019

No. 135 Rev. G

Signature: 