


# ActiGraph™

## Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

MANUFACTURER		
Name of Company	Address	SRN
ActiGraph L.L.C.	70 North Baylen Street, Suite 400, Pensacola, FL. 32502 United States of America	US-MF-000007954

AUTHORIZED REPRESENTATIVE			
Name of Company	Address	SRN	Telephone/fax/email
Emergo Europe	Westervoortsedijk 60 6827 AT Arnhem The Netherlands	NL-AR-000000116	+31.70.345.8570 - phone EmergoEurope@ul.com

PRODUCT IDENTIFICATION		
Product Name	Product Code / Catalog Number	Basic UDI-DI
ActiGraph LEAP™	LEAP	00853048008053
Intended Purpose	Photo	
The ActiGraph LEAP™ is a small, worn activity monitor designed for documenting physical movement associated with applications in physiological monitoring. The device is intended to monitor the activity associated with movement during sleep. The ActiGraph LEAP™ can be used to analyze circadian rhythms and assess activity in any instance where quantifiable analysis of physical motion is desirable.		

RISK CLASS FOR MEDICAL DEVICES	
Device Classification	Common Specifications
Class: I, Self-Certified (active, non-measuring) Rules: 13 (Annex VIII)	EN 55032: 2015 + A11:2020 + A1:2020 EN 61000-3-3:2013 + A1:2019 + A2:2021 EN IEC 61000-3-2:2019 + A1:2021 EN 55035:2017 + A11:2020 EN 301 489-1 V2.2.3 EN 301 489-17 V3.2.4 EN 300 328 V2.2.2 EN 62479:2010



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	EN IEC 62368-1:2020 + A11:2020 IEC 60601-1:2005+A1:2012+A2:2020; EN 60601-1:2006+A1:2013+A2:2021 Basic Safety and Essential Performance IEC 60601-1-2:2014; EN 60601-1-2:2015 EMC IEC 60601-1-6:2010 + A1:2013; EN 60601-1-6:2010+A1:2015 Usability IEC 60601-1-9:2007+A1:2013; EN 60601-1-9:2008+A1:2013 Environmentally Conscious Design IEC 60601-1-11; EN 60601-1-11:2015+A1:2021 Home Healthcare
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ActiGraph declares that the above-mentioned products meet the provision of the following EU legislation:

- Medical Device Regulation, MDR (EU) 2017/745
- Radio Equipment Directive (RED) 2014/53/EU
- Restriction of the use of certain hazardous substances (RoHS) Directive 2011/65/EU

**COMPANY REPRESENTATIVE:** Thomas Hartshorn

**TITLE:** Head of Quality Assurance & Compliance

**PLACE:** Pensacola, FL. United States of America

**SIGNATURE:**

**DATE:**

22 Mar 2024