

Declaration of Conformity - PRO-Diary

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

MANUFACTURER			AUTHORISED REPRESENTATIVE	
Name of Company	Address	Nar	me of Company	Address
CamNtech Limited	Riband House Manor Farm Fenstanton	Qar	ad EC-REP BV	Pas 257 2440 Geel Belgium
SRN: GB-MF-000004683	Cambridgeshire PE28 9JD, UK	geshire SRN	1: BE-AR-000000040	
	PRODUCT II	DENTIF	ICATION	
Product Name	Catalogue	No.	Basic LIDI-DI	

PRODUCT IDENTIFICATION				
Product Name	Catalogue No.	Basic UDI-DI		
PRO-Diary	05-111	++B329PDM01XT		
PRO-Diary V	05-121			

Intended Purpose Representative Photos

The PRO-Diary is a compact, lightweight, body-worn activity monitoring device that may be used to document physical movement associated with applications in physiological monitoring. The device is intended to monitor limb or body movements during daily living and sleep. The PRO-Diary can be used to assess activity in any instance where quantifiable analysis of physical motion is desired.



RISK CLASS FOR MEDICAL DEVICES

Class 1 (active, non-measuring, non-sterile) Annex VIII of regulation 2017/745; rules 1, 13 Conformity Assessment: Annex II and Annex III of regulation 2017/745

CamNtech declares that the above-mentioned products meet the provision of the following EU legislation:

- Medical Device Regulation (MDR) 2017/745
- RoHS Directive 2011/65/EU
- WEEE Directive 2012/19/EU

COMPANY REPRESENTATIVE: Howard Smith

TITLE: Director SIGNATURE:

PLACE: United Kingdom **DATE:** 21/05/2021