

## **EU Declaration of Conformity**

## Manufacturer:

iTs Designs Ltd. (T/A: Alert-iT Care Alarms) Fernie House (Unit 3) Coalville Business Park Coalville, Leicestershire LE67 3NR, England

Tel: +44 (0) 1530 239 900

Single Registration Number (SRN) - XXXXXXXX

## **European Authorised Representative:**

LEISNER ApS Korsvangcentret DK-5610 Assens Denmark

Tel: +45 6371 3050

Model:	Description:	Basic UDI-DI (GMN):	UDI-DI (GTIN-14):
P176AAA	Companion mini: 434 MHz, Branding	506053070P176AALE	50605307017734
P176ABA	Companion mini: 869 MHz, Branding	506053070P176ABLG	50605307017802
P176AAB	Companion mini: 434 MHz, Branding	506053070P176AALE	50605307017970
P176ABB	Companion mini: 869 MHz, Branding	506053070P176ABLG	50605307018038
P176AAC	Companion mini: 434 MHz, Branding	506053070P176AALE	50605307018106
P176ABC	Companion mini: 869 MHz, Branding	506053070P176ABLG	50605307018274
P176AAG	Companion mini: 434 MHz, Branding	506053070P176AALE	50605307018342
P176ABG	Companion mini: 869 MHz, Branding	506053070P176ABLG	50605307018410

Description:	The "Companion mini" detects movements and sends a failsafe alarm message over Safelink <sup>™</sup> radio if the movements exceed user defined limits.	
Directives:	2017/745	Medical Device Regulation (MDR)
2	2014/53/EU	Radio Equipment Directive (RED)
	2011/65/EU + 2015/863	Permitted Materials (RoHS)
	2012/19/EU	Waste Electrical and Electronic Equipment Directive (WEEE)
Standards Applied:	BS EN 60601-1:2006+A12:2014	Medical Equipment: Basic Safety Requirements
	BS EN 60601-1-2:2015+A1:2021	Medical Equipment: EMC Requirements
	BS EN 60601-1-6:2010+A1:2015	Medical Equipment: Usability Requirements
	BS EN 60601-1-8:2007+A11:2017	Medical Equipment: Alarm Systems
	BS EN 60601-1-11:2015	Medical Equipment: Home Healthcare Requirements
	EN 300-220-2 v3.1.1	Radio Emissions (P176AA 434MHz Only)
	EN 300-220-3-1 v2.1.1	Radio Emissions (P176AB 869MHz Only)

We hereby, under our sole responsibility declare, that the product listed above, is in conformity with the Medical Device Regulation (MDR) 2017/745 as a class I medical device based on Annex 8 and the relevant European harmonised standards listed above.

The product concerned has been manufactured under a quality management system (ISO 13485:2016) according to Annex 9 of MDR 2017/745 EU and audited by BSI.

This EU declaration of conformity was written accordance to Annex 4 of the MDR, and all supporting documentation is retained at the premises of the manufacturer.

Date: OI JUNE

Place: Leicester, UK.

Name: RICHARD GUNN Position: DIRECTOR

2021