



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™ radio if the movements exceed user defined limits.	
Directives:	2017/745 2014/53/EU 2011/65/EU + 2015/863 2012/19/EU	Medical Device Regulation (MDR) Radio Equipment Directive (RED) Permitted Materials (RoHS) Waste Electrical and Electronic Equipment Directive (WEEE)
Standards Applied:	BS EN 60601-1:2006+A12:2014 BS EN 60601-1-2:2015+A1:2021 BS EN 60601-1-6:2010+A1:2015 BS EN 60601-1-8:2007+A11:2017 BS EN 60601-1-11:2015 EN 300-220-2 v3.1.1 EN 300-220-3-1 v2.1.1	Medical Equipment: Basic Safety Requirements Medical Equipment: EMC Requirements Medical Equipment: Usability Requirements Medical Equipment: Alarm Systems Medical Equipment: Home Healthcare Requirements Radio Emissions (P176AA 434MHz Only) 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: 01 JUNE 2021

Place: Leicester, UK.

Name: RICHARD GUNN

Position: DIRECTOR