

EC Declaration of Conformity

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

MANUFACTURER			
Name of Company	Address	SRN	
TechTrade LLC	6900 Tavistock Lakes Blvd.	US-MF-000023808	
	Suite 400		
	Orlando, FL 32827 USA		

AUTHORIZED REPRESENTATIVE			
Name of Company	Address	SRN	Phone/email
Emergo Europe	Westervoortsedijk 60	NL-AR-00000116	+31.70.345.8570
	6827 AT Arnhem		EmergoEurope@ul.com
	The Netherlands		

PRODUCT IDENTIFICATION Product Name		Code / Catalog Number
Ready-Heat One Panel Blanket Ready-Heat Four Panel Blanket Ready-Heat Six Panel Blanket Ready-Heat Vest Ready-Heat Infant Warming Blank Ready-Heat II Blanket Ready-Heat Temperature Manage Ready-Heat Infant Warming Matti Ready-Heat II Torso Blanket Ready-Heat Temperature Manage	ement Full Body ress	S1RHSM S4RHMD/S4RHMD-N S6RHLG GRHV06A GIW6C G12RH2 SB9RH9120 S2RHIM G6RH2T/G6RH2TB SB6RH9120
Ready-Heat Emergency and Disaster Blanket		ED9RH
Intended Purpose		Basic UDI-DI
To provide warm relief, aid in the prev	vention of hypothermia	0850017905100SB
CDN/EMDN Description and Code:	Body Thermoregulation Equipme	ent – Z12040208

RISK CLASS F	OR DEVICES		
Device Classifi	cation	Common Specifications / Standards	Technical File Information
Class:	lla		TF-01 Rev Q Dec 2023
Rule:	9		

NOTIFIED BODY			
Name of Company	ID Number	Conformity Assessment Procedure	Certificate Reference(s)
GMED SAS	CE0495	9	Cert. 36083 rev.2, Add.
		MDD 93/42/EEC Council	38580 rev.5, Conf.
		Directive	Letter 39487 rev.0



TechTrade LLC., hereby declares under our exclusive responsibility the above-mentioned products meet the relevant provisions of the European Regulation (EU) 2017/745 for Medical Devices and those General Safety and Performance Requirements listed in Annex I; also, any applicable standards, any Common Specifications, or related European Union legislation. The conformity of the device is confirmed through placement of the CE Mark on each device. All supporting documentation is retained under the premises of the manufacturer.

List of Harmonized Standards, Common Specifications, and other relevant EU legislation

Identification Number	Title or Description	Version or Year
ISO 13485	Medical devices — Quality management systems — Requirements for regulatory purposes	2016
MDR EU 2017- 745	European Regulation (EU) 2017/745 for Medical Devices	2017
EN ISO 14971 +A11	Medical devices — Application of risk management to medical devices	2019 2021
EN 62366-1 + AMD1	Medical devices. Application of usability engineering to medical devices	2015 2020
EN ISO 15223-1	Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements	2021
ISO 10993-1	Biological evaluation of medical devices Part 1: Evaluation and testing within a risk management process	2020
EU 207/2012	Regulation (EU) No 207/2012 on electronic instructions for use of medical devices	2012
ISO 20417	Information Supplied by the Manufacturer.	2021

COMPANY REPRESENTATIVE: Erin Bart

TITLE: Regulatory Manager SIGNATURE:

PLACE: Orlando, FL, USA DATE: EU 31/01/2024