

## **EC CERTIFICATION**

# FULL QUALITY ASSURANCE SYSTEM

## Directive 93/42/EEC on Medical Devices, Annex II excluding (4)

We hereby declare that an examination of the under mentioned full quality assurance system has been carried out following the requirements of the Swedish national legislation LVFS 2003:11 to which the undersigned is subjected, transposing Annex II (with the exemption of section 4) of the Directive 93/42/EEC on medical devices. We certify that the full quality assurance system conforms with the relevant provisions of the aforementioned directive, and the result entitles the organization to use the CE 0413 marking on those products listed below.

#### Organization:

## **Kanmed AB**

Main Site: Gårdsfogdevägen 18B, SE-168 67 Bromma, Sweden

#### **Product Category:**

- Patient Warming Systems

For further identification of the products covered, see the MDD product list/product schedule.

#### **Certificate Number:**

41313430-05

#### **Initial Certification Date:**

19 June 2000

#### Certificate Valid from:

9 March 2021

#### **Certificate Expiry Date:**

26 May 2024



#### **Peter Nermander**

Certification Authority MDD Intertek Semko AB, Kista, Sweden

9 March 2021

#### **Signed Date**

Intertek Semko AB Box 1103, SE-164 22 Kista, Sweden Telephone +46 8 750 00 00 medtechsweden@intertek.com

The certification is subject to the organization maintaining their system in compliance with the regulations stated in this certificate, allowing regular assessments and following the contracted requirements of the Notified Body.

Intertek Semko AB is a Notified Body according to Directive 93/42/EEC on medical devices, with identification number 0413.







### MDD - Product List

Products included in the Certificate No:

Issued to:

41313430-05

Kanmed AB

Gårdsfogdevägen 18B SE-168 67 Bromma

Sweden

Product category	Type/Model designation	Class	Sterile	GMDN code (not mandatory)	Date added
Patient Warm	ing Systems				
	Patient Warming				
	Heating Pads				
,	OP3-063, 63 cm	llb	=	-	*
	OP3-104, 104 cm	llb	-		*
	OP3-150, 150 cm	IIb	-		*
	WarmCloud				
	9000	IIb	No	37328	Feb 03, 2017
	Infant Warming				
	Baby Warming System				***
	BW3-020	IIb	-		March 15, 2011
	Heating Pads				
	BW-50-002	IIb	=		*
	BW3-003	IIb			March 15, 2011

<sup>\*</sup> Product added before June 19, 2010.

Date of Issue: 9 March 2021

Intertek Semko AB **Notified Body MDD** 

Peter Nermander Certification Authority MDD

This product list is only valid together with the referenced, valid EC certificate.

The GMDN codes are assigned by the manufacturer and are only provided for convenience.

Intertek Semko AB is a Notified Body according to the Directive 93/42/EEC on medical devices, with identification number 0413.

Product List for Certificate No: 41313430-05

Date: 9 March 2021



## MDD - Decision Report

Certificate No:

41313430-05

Date:

9 March 2021

Handled by: E-mail: medtechsweden@intertek.com

Nina Fazil

Kanmed AB

Attn: Annika Engstrand Gårdsfogdevägen 18B SE-168 67 Bromma Sweden

**Purpose** 

Assessment of notification received from client regarding a correction of

the company name from KanMed AB to Kanmed AB according to the national legislation for medical devices LVFS 2003:11 (Medical Device

Directive 93/42/EEC), Annex II.

**Activity** 

The correction has been assessed and accepted.

Additionally the address format was corrected (post code and city were

swapped).

Scope of assessment

- Patient Warming Systems, Class IIb

Certificate Valid from

9 March 2021

Conclusions/Decisions

Referring to the above a Certificate of Conformance with the national legislation for medical devices LVFS 2003:11 (Medical Device Directive 93/42/EEC), Annex II will be reissued with the correct company name. The Certificate is valid for products specified in the "MDD - Product List".

**Appeals** 

Any appeal against this decision will be processed by an appeals panel as

Intertek. The appeal shall be submitted to Intertek Semko AB, PO-Box

1103, SE-164 22 Kista, Sweden.

**Others** 

Any complaints, from customers and others, and corrective actions concerning your certified quality system shall be documented and retained. Upon request Intertek Semko has the right to review this

documentation.

Intertek Semko AB Notified Body MDD

Peter Nermander

Certification Authority MDD