Declaration of Conformity to Directive 93/42/EEC concerning Medical Devices FORM 140-05 / DCR 13-459
Document: RA DoC-020 Rev. 10 / DCR 19-100
Climber Guiding Catheter

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DECLARATION OF CONFORMITY to Directive 93/42/EEC concerning Medical Devices

Name of Product: Climber™ Guiding Catheter

Legal (labelled) PendraCare International B.V.

Manufacturer: Van der Waalspark 22

9351 VC Leek The Netherlands

Dutch Chamber of Commerce - Registration Number 02086018

Declaration:

I hereby declare that the medical device specified in this declaration conforms to the provisions of the *current* European Council (EC) Directive 93/42/EEC of June 14, 1993 concerning Medical Devices and therefore bears the CE mark of conformity on its labelling in combination with the Notified Body Identification number 0344 of DEKRA Certification B.V., Arnhem, The Netherlands.

- The product conforms to the applicable Essential Requirements for Safety and Performance per current Directive 93/42/EEC, Annex I: "Essential Requirements".
- The device classification (i.e., Class III) has been determined per *current* Directive 93/42/EEC, Annex IX: Classification Criteria,
- The appropriate Conformity Assessment module per article 11 of the *current* Directive 93/42/EEC, i.e., Annex II (4) has been followed as indicated on the "EC Design-Examination" Certificate (2020764DE02) in combination with this Declaration of Conformity,
- PendraCare's Quality Management System (QMS) fulfils the requirements described in the *current* Directive 93/42/EEC and EN-ISO 13485: 2016 as evidenced by the "CE Marking of Conformity" Certificate [2020764CE01- Full Quality Assurance System per Annex II excluding (4)] and its accompanying Certification Notice (2020764CN) and by the Certificate of Registration (2086817). The specified medical device falls within the scope of PendraCare's QMS as indicated in the certificates.

GMDN: GMDN Term*: Vascular Guide Catheter GMDN Code*: 17846

* per GMDN agency database

Valid until: This Declaration of Conformity is valid until **November 1, 2023**, i.e., the validity date

indicated on the CE-marking of Conformity Certificate and the product's EC Design-

Examination Certificate.

Reference: RA DoCa-020 Rev. 10 - Annex to the Declaration of Conformity.

Place of issue: Leek, The Netherlands

Date: 2019-03-26

Declared by:

J. van der Kuil

Manager Quality Assurance & Regulatory Affairs

