



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

Name of Product: **Alviguide™ Blue+ Interventional Cardiology Guiding Catheter**

Legal (labeled) Manufacturer: **Alvimedica Tibbi Ürünler Sanayi ve Dış Ticaret A.Ş.**
İstanbul Trakya Serbest Bölgesi Ferhatpaşa Mah. Atatürk Bulvarı Manolya Sk.
No: 7 Çatalca 34540, İstanbul, Turkey

Declaration:

We, the undersigned, 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.**

- Conformity to the applicable Essential Requirements for Safety and Performance per *current* Directive 93/42/EEC, Annex I: "Essential Requirements" has been proven,
- 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, Section 4**) has been followed as indicated on the "EC Design Examination" Certificate (**2161507DE04**) in combination with this Declaration of Conformity,
- Alvimedica's Quality Management System fulfils the Quality Management System requirements described in the *current* Directive 93/42/EEC (**Annex II, excluding Section 4**) and **EN ISO 13485: 2012** as evidenced by the "CE Marking of Conformity" Certificate (**2161507CE02**), its accompanying Certification Notice and the Certificate of Registration (**2161507**). The specified medical device falls within the scope of Alvimedica's Quality Management System as indicated in the Certificates.

GMDN: GMDN Term*: **Intravascular Guiding Catheter** GMDN Code*: **17846**
* per GMDN agency database

Valid until: This Declaration of Conformity is valid until **June 1, 2018**, i.e. the validity date indicated on the "CE Marking of Conformity" Certificate issued by DEKRA.

Reference: RA-DoCa-002 Rev. **014** - Annex to the Declaration of Conformity.

Place of issue: **İstanbul, Turkey**

Declared by: **Nursen Erin** Date: **2017-08-07**

Quality Assurance Manager