



## EU DECLARATION OF CONFORMITY

Manufacturer: Duckworth & Kent Ltd  
Terence House  
7 Marquis Business Centre  
Royston Road  
Baldock, Hertfordshire  
SG7 6XL, United Kingdom

Authorised Representative: Emergo Europe  
Prinsessegracht 20  
2514 AP The Hague  
The Netherlands

Duckworth & Kent Ltd declares that the medical devices listed in Table 1 and described hereafter,

### **Duckworth & Kent range of Invasive Diamond Knives**

are in conformity with the essential requirements and provisions of the Council Directive 93/42/EEC as amended by Directive 2007/47/EC and as transposed into UK legislation (UK Medical Devices Regulations SI 2002 No. 618, as amended).

In accordance with Annex VII of the Council Directive 93/42/EEC as amended by Directive 2007/47/EC and as transposed into UK legislation (UK Medical Devices Regulations SI 2002 No. 618, as amended),

### **These devices are designated as Class I**

#### Classification Route

These devices are all transient use, reusable, non-sterile, invasive medical devices.

The classification is determined under Annex IX, Rule 5

This declaration is issued under the sole responsibility of Duckworth & Kent Ltd.

Martin Lock  
Head of Regulatory Affairs



**Table 1. Medical Devices covered under this Declaration of Conformity.**

Diamond Knives, Invasive	
REF	Device Title
4-100	Diamond Knife
4-115	Diamond Knife
4-122	Diamond Knife
4-125	Diamond Knife
4-402	Diamond Knife
4-406	Diamond Knife
4-415	Diamond Knife
4-416	Diamond Knife
4-430	Diamond Knife
4-438	Diamond Knife
4-440	Diamond Knife
4-460	Diamond Knife
4-480	Diamond Knife
4-590	Diamond Knife
4-600	Diamond Knife
4-620	Diamond Knife
4-620-3	Diamond Knife
4-620-4	Diamond Knife
4-621	Diamond Knife
5-300-1	Diamond Knife
5-305-1	Diamond Knife
5-310-1	Diamond Knife
5-329-1	Diamond Knife
5-330-1	Diamond Knife
5-340-1	Diamond Knife
5-360-1	Diamond Knife
5-362	Diamond Knife
6-607	Lamellar Dissector



**Table 2. Relevant standards and common specifications.**

Applicable Standards	
Standard Number	Title
EN 1041:2008	Information supplied by the manufacturer of medical devices
EN ISO 10993-1:2009	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process (ISO 10993-1:2009) EN ISO 10993-1:2009/AC:2010
EN ISO 13485:2016	Medical devices - Quality management systems - Requirements for regulatory purposes (ISO 13485:2016) EN ISO 13485:2016/AC:2018
EN ISO 14971:2012	Medical Devices - Application of risk management to medical devices (ISO 14971:2007, Corrected version 2007-10-01)
EN ISO 15223-1:2016	Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements (ISO 15223-1:2016, Corrected version 2017-03)
EN ISO 17664:2017	Processing of health care products - Information to be provided by the medical device manufacturer for the processing of medical devices (ISO 17664:2017)
EN 62366-1:2015+A1:2020	Medical devices - Part 1: Application of usability engineering to medical devices
93/42/EEC	Council Directive concerning medical devices
2007/47/EC	Council Directive amending 93/42/EEC
SI 2002 No. 618	Medical Devices Regulations (UK)
MEDDEV 2.7/1 rev.4 June 2016	Clinical evaluation: A guide for manufacturers and notified bodies
MEDDEV 2.12/1 rev.8 Jan 2013	Guidelines on a medical device's vigilance system



## ADDENDUM TO THE ORIGINAL DECLARATION OF CONFORMITY

Per 31 January 2023, the address of our EU Authorised Representative as listed on the original Declaration of Conformity has changed

Old Address  
Authorised Representative: Emergo Europe  
Prinsessegracht 20  
2514 AP The Hague  
The Netherlands

New Address  
Authorised Representative: Emergo Europe  
Westervoortsedijk 60  
6827 AT Arnhem  
The Netherlands

Martin Lock  
Head of Regulatory Affairs

25/01/23

Date