schülke -}

EC declaration of conformity

according to Annex II - excluding Section 4 of Council Directive 93/42/EEC concerning medical devices

Medical Device	rotasept [®]
Manufacturer	Schülke & Mayr GmbH Robert-Koch-Str. 2 22851 Norderstedt
Notified Body	DQS Medizinprodukte GmbH August-Schanz-Str. 21 60433 Frankfurt am Main Ident.No.: 0297
Classification acc. to Directive 93/42/EEC, Annex IX, Rule 15	Ilb
Product group	Disinfectant, dental
Product category	05 - Hospital hardware
Issued CE certificates by DQS Medizinprodukte GmbH/0297	EN ISO 9001 – Cert. Reg. No. 004567 QM08 EN ISO 13485 – Cert. Reg. No. 004567 MP2012 Annex II – Cert. Reg. No. 004567 MR2
Standards applied	Applied standards are listed in Sec. 2.4 of the technical documentation. Location of technical documentation: Schülke & Mayr GmbH, Reg. Affairs

We herewith declare that the described device corresponds to the essential requirements of the EEC directive concerning medical devices.

l, the undersigned, declare that Schülke bears the sole responsibility for issuing this Declaration

Norderstedt, 17.12.2015

ppa. Dr. W. Weltger

Head of Quality Management

Schülke & Mayr GmbH

ppa Dr. P. Oltmanns

Head of Research & Development

Muning

Schülke & Mayr GmbH

This Declaration is valid until an updated version has been issued, but not longer than 2018-10-17.