



EC declaration of conformity

Medical Device name	perform®		
Formulation No.	F08		
Product group	Disinfectant, medical device surfaces		
Product Category	05 - Hospital hardware		
Intended Purpose	surface disinfectant		
Risk Class	II a		
according to Directive 93/42/EEC	annex	IX	
Standards applied	EN ISO 13485 additional standards see technical documentation Schülke & Mayr GmbH, Regulatory Affairs		
Manufacturer	Schülke & Mayr GmbH		
according to Directive 93/42/EEC	Robert-Koch-Str. 2 22851 Norderstedt Germany		
Notified Body	DQS Medizinprodukte GmbH August-Schanz-Str. 21 60433 Frankfurt am Main Germany Ident.No.: 0297		
Conformity Assessment Procedure	Annex II excluding section 4		
according to Council Directive 93/42/EEC			
Issued Certificates	Annex II 93/42/EEC	Cert. Reg. No.	004567 MR2
Version	9.0		

Schülke & Mayr GmbH herewith declares that the device covered by this declaration is in conformity with the Council Directive 93/42/EEC concerning medical devices.

Schülke & Mayr GmbH declares that Schülke & Mayr GmbH bears the sole responsibility for issuing this Declaration

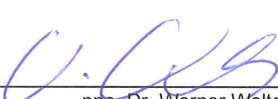
Norderstedt

06.05.2019

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