



## EU Declaration of Conformity

|                                                                     |                                                                                                      |       |                                                                     |
|---------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|-------|---------------------------------------------------------------------|
| Manufacturer<br>according to Regulation 2017/745                    | Schülke & Mayr GmbH<br>Robert-Koch-Str. 2<br>22851 Norderstedt<br>Germany                            |       |                                                                     |
| Registration Number<br>acc. to Art. 31 2017/745                     | DE-MF-000005701                                                                                      |       |                                                                     |
| <b>Product Name</b>                                                 | <b>gigasept® Instru AF</b>                                                                           |       |                                                                     |
| Basic UDI-DI<br>Code acc. to Art. 26 2017/745                       | 4032651BSC00000037AH<br>Z12011385                                                                    |       |                                                                     |
| Intended Purpose                                                    | cleaning and disinfection agent for manual reprocessing of medical devices                           |       |                                                                     |
| Risk Class<br>according to Regulation 2017/745                      | II a                                                                                                 | Annex | VIII                                                                |
|                                                                     |                                                                                                      | rule  | 16                                                                  |
| Standards applied                                                   | EN ISO 13485<br>additional standards see technical documentation<br>Schülke & Mayr GmbH              |       |                                                                     |
| Notified Body                                                       | DQS Medizinprodukte GmbH<br>August-Schanz-Str. 21<br>60433 Frankfurt am Main<br>Germany<br>No.: 0297 |       |                                                                     |
| Conformity Assessment Procedure<br>according to Regulation 2017/745 | Annex                                                                                                | IX    | Chapter I, II section 4 and III                                     |
| Certificates                                                        | Annex                                                                                                | IX    | 004567 MDR2017Q<br>004567 MDR2017B<br>EN ISO 13485<br>004567 MP2016 |
| Version                                                             | 1-0                                                                                                  |       |                                                                     |

Schülke & Mayr GmbH herewith declares that the device covered by this declaration is in conformity with the Regulation 2017/745 concerning medical devices.

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

Norderstedt

15.06.2023  
ppa.

  
Dr. Sven Pfleging  
Schülke & Mayr GmbH  
Chief Commercial Officer

15.06.2023  
ppa.

  
Lars Lemke  
Schülke & Mayr GmbH  
Chief Operating Officer