

EC Declaration of Conformity

in Accordance with EC Directive 93/42/EEC on Medical Devices

Manufacturer Carl Zeiss Meditec AG, Goeschwitzer Strasse 51-52, 07745 Jena, Germany

We Carl Zeiss Meditec AG herewith declare with sole responsibility that the following Medical Device meets the Requirements of the European Directive 93/42/EEC. The device is provided with CE Marking.

Product identification:

Intraocular lens

Medical Device Trade Name:

AT TORBI 709M

Models/Reference:

Accessories:

Including all options and accessories as defined in the

User Manual

Medical Device Class

Class IIb

MDD 93/42/EEC:

Conformity Assessment Procedure

Annex II of MDD 93/42/EEC

Scope of Application:

This Declaration of Conformity is valid for all products

manufactured until 2021-11-29

UMDNS classification:

16-071

GMDN Code

35658

Notified Body: DQS Medizinprodukte GmbH, August-Schanz-Straße 21, 60433 Frankfurt – notified under 0297. Any Modification to the Product not authorized by Carl Zeiss Meditec AG will invalidate this Declaration.

i.V. Guillaume Gasc

Vice President Competence Center &

Order fulfillment consumables

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Implants / OVDs / Sterile Products

Berlin, 2016-11-22

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