

EU DECLARATION OF CONFORMITY According to Annex II and Annex III of Regulation (EU) 2017/745

GC EUROPE N.V. Research Park Interleuvenlaan 33 B-3001 Leuven Belgium

SRN: BE-MF-000001608

We ensure and declare under our sole responsibility that the product :

Product Name	Classification according to Annex VIII of the Regulation (EU) 2017/745	Article List	Basic UDI-DI
GC Fuji VARNISH	Class I, Rule 5	According to the Attachment	++J022MD0002JN

to which this declaration relates is in conformity with the following standards or other normative documents :

EN ISO 13485:2016 Medical Devices –Quality Management Systems – Requirements for Regulatory Purposes

and meets the provisions of Regulation (EU) 2017/745 concerning Medical Devices which apply to it, and is manufactured in accordance with the technical documentation.

Leuven, 10 May 2021 Date

Mario Minale Head of Regulatory Affairs On behalf of GC EUROPE N.V.





LIST OF PRODUCTS

Basic UDI-DI		New Article code	Description
++J022MD0002JN	000026	10000002	GC Fuji VARNISH, 10.4ml (10g) Bottle

