



# Declaration of Conformity

Manufacturer                      CuraMedical B.V.  
Address                              Industrieweg 6B  
   1566 JP ASSENDELFT  
   The Netherlands

Product Type                      Oxidized Regenerated Cellulose Hemostat, Sterile, Absorbable, Single Use  
Classification                      III  
(MDD, Annex IX)

GMDN code                         38771

Intended use                        To be used in surgical procedures to control capillary, venous and small  
arterial bleeding when binding or other conventional methods of control are  
impractical or ineffective.

Brand name                         **CuraCel® High Density**

Dye                                      Un-dyed (natural, beige-caramel)

Insulation / Coating              NA

Types and Sizes:

CH-603	26 x 26 mm	CH-637	76 x 102 mm
CH-607	50 x 75 mm	CH-660	152 x 230 mm

*We herewith declare under our sole responsibility that the above mentioned products meet the provisions of the following EC Council Directives and Standards. The design of the products and the Quality System meets the requirements according to Annex II – Section 3 and 4 of the Council Directive 93/42/EEC of June 14<sup>th</sup>, 1993 concerning medical devices. All supporting documentations are retained under the premises of the manufacturer and the notified body.*

## DIRECTIVES

### General applicable directives:

Medical Device Directive: COUNCIL DIRECTIVE 93/42/EEC of June 14<sup>th</sup>, 1993 concerning medical devices (MDD 93/42/EEC), as last amended September 5<sup>th</sup>, 2007 (Directive 2007/47/EC).

### Standards:

All relevant Harmonized Standards as published in the Official Journal of the European Communities and listed in the latest version of document: QS-1530-01 List of applicable standards Curacel, are applicable to this type of product.

United States Pharmacopoeia, USP Monographs **OXIDIZED REGENERATED CELLULOSE**.

Notified Body                        MDC Medical Device Certification GmbH,  
   Kriegerstrasse 6, D-70191 Stuttgart, Germany  
   Notified Body number: 0483

Certificate no.                        D1169400018                      Expiry Date    2023-01-09

Date CE mark was first affixed    10-02-2010

Place & Date                         ASSENDELFT, 25-06-2018

Signature

Name

F. J. Hoogland

Position

Director Marketing & Sales and Regulatory Affairs

