SD BIOSENSOR

Declaration of Conformity

Manufacturer Name SD Biosensor, Inc. Manufacturer Address Head Office C-4th&5th, 16, Deogyeong-daero, 1556beon-gil, Yeongtong-gu, Suwon-si, Gyeonggi-do 16690, KOREA Manufacturing Site 74, Osongsaengmyeong 4-ro, Osong-eup, Heungdeok-gu, Cheongju-si, Chungcheongbuk-do 28161, KOREA **MT Promedt Consulting GmbH EC Representative Name EC Representative Address** Altenhofstrasse 80 66386 St. Ingbert Germany **Common Name** Immunoassay Test Kit STANDARDTM F hs-CRP **Product Name** STANDARDTM F hs-CRP Control *Please refer to "Annex I. Product List" on page 2 in more detail. **Reference Number** F-HCRP, C-F-HCRP Classification Others not covered by Annex II and self-testing according to Directive 98/79/EC **Conformity Assessment Route** Annex III of IVDD 98/79/EC (EC Declaration of Conformity) EN ISO 13485:2016 EN ISO 18113-1:2011 **Applied Standards** EN ISO 14971:2012 EN ISO 18113-2:2011 EN ISO 23640:2015 EN ISO 15223-1:2016 EN ISO 17511:2003 EN 62366:2008 EN 13612:2002

We herewith declare that the above mentioned products meet the provisions of the Council Directive 98/79/EC for in vitro diagnostic medical devices. All supporting documentations are retained under the premises of the manufacturer. SD Biosensor, Inc. is exclusively responsible for the declaration of conformity.

Place: Suwon-si, Republic of Korea Valid from: May 13, 2022

Signature

Hyo-Keun, Lee CEO / President

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Annex I. Product List

 F-HCRP STANDARD[™] F hs-CRP Test Device	EDMA Code	Description of EDMA code
(individually in a foil pouch with desiccant and Spoit [™]) Extraction buffer Instructions for use	12 11 01 09 00	C-Reactive Protein
 <u>C-F-HCRP</u> STANDARD™ F hs-CRP Control STANDARD F hs-CRP Control – Level 1 (10 tablets in the bottle, Transparent tube) STANDARD F hs-CRP Control – Level 2 (10 tablets in the bottle, Red colored tube) Instructions for use 	EDMA Code 12 50 01 06 00	Description of EDMA code Specific Protein Controls