CERTIFICATE

With EU Directive of MDD 93/42/EEC

Certificate No.:

CE-19-0822-01

Applicant:

Longkou Sanyi Medical Equipment Co., Ltd.

West Rost, 2 Klometers North Of Huangcheng

Bus Station, Longkou, Shandong

Product:

Medical X-ray Protective Equipment and Device

Model(s):

FA01/FA01-2/FA02/FA02-2/FA03/FA03-2/FA04/FA05/FA05-2/FA06/ FA07/FA08/FA09/FA10/FA11/FA12/FA13/FA14/FA15/FA16/FA17/ FAA01/FAA02/FAA03/FAA04/FAA05/FAA06/FAA07/FAA08/ FAA10/FAA11/FAA15/FC01/FC02/FC03/FC04/FC05/FC06/FC07/ FC08/FC09/FC10/FC11/FC12/FC13/FC14/FC15/FC16/FC17/FC18/ FC19/FC20/FC21/FE01/FE02/FE03/FE04/FE05/FE06/FE07/FE08/ FE09/FE09-1/FE10/FE11/FW01/FW02/FW03/FW04/FW05/FW06/ FW07/FW08/FW09/FW09-1/FW10/FW11/FG01/FG02/FG03/FG04/ FG05/FG06/FG07/FG08/FG09/FG10/FG11/FG12/FG13/FG14/FG15/

FG16/FG17/FG18/FG19/FG20/FG21/FG22/FG23/FG24/FG25/FG26

Standards:

EN ISO 14971:2012,EN ISO 15223-1:2016

EN 1041:2008, EN ISO 10993-1:2009/AC:2010

Based upon the voluntary assessment of the product sample and Technical Construction File, the apparatus is deemed to meet the requirements of the above standards and EC directives.

The manufacturer has the responsibility for ensuring that all serial manufacture of the products are in compliance with the specification of the sample submitted for assessment and detailed in the technical file.

CE

Date: 22/08/2019

Stamp:

Solutions GmbH

The CE marking may be used if all relevant and effective EC directives are complied with.

Euroscene Business Solutions Limited

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