

# *EC Declaration of Conformity*

MANUFACTURER  
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EC REP  
**S.B. PHARMA GmbH**

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**Proposed Scope :**

**93/42/EEC–(Annex II  
(excluding section 4)) –  
CE1639**

Certificate K21/81826527

**93/42/EEC – (Annex V) –  
CE1639**

Certificate K21/81826528

**ISO 13485:2016 – UKAS**

Certificate KR20/81826471

**Infrared forehead thermometer Model: FS-700, HFS-700, HFS-1000  
Nasal Aspirators Model:HNA-200, HNA-300, HNA-1000**

**Blood pressure monitor Model:HBP-500, HBP-700, HBP-1520**

**Design, development and manufacture of infrared thermometers, blood pressure  
monitor and nasal aspirators**

The manufacturer is exclusively responsible for declaration of conformity.

Declares that the medical device described hereafter,

**Product name:** Infrared Forehead Thermometer  
**Model Name:** FS-700  
**UMDNS CODE:** 14036 [Thermometers, Infrared]  
**Classification:** Class II a(Annex IX Rule10)

Noted product is in conformity with technical requirements and applicable regulations:

**Directive:** 93/42/EEC amended by MDD 07/47/EC

**Standards:** EN 60601-1:2006  
EN 60601-1-6:2010  
EN 60601-1-11:2010  
ISO 80601-2-56:2017  
EN 60601-1-2:2015  
EN 1041:2008  
EN ISO 15223-1:2016

**Notified Body:** SGS Belgium NV  
Noorderlaan 87, BE-2030 Antwerpen, Belgium  
Notified Body Number 1639

This product is classified as Class II a, Council Directive 93/42/EEC, Annex IX, Rule 10, and complies with the requirements and regulations of the Council Directive 93/42/EEC as amended by Directive 2007/47/EC.

**Date of issued: March 9<sup>th</sup> 2021**



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**JAE-HO SHIN**

**PRESIDENT**

**on behalf of HuBDIC Co.,Ltd.**