

## EC DECLARATION OF CONFORMITY

## According to Annex II (exemption of section 4) of Council Directive 93/42/EEC concerning Medical Devices

Reference No

1828C-11

Manufacturer

SIDAPHARM P.C.

21, Stageiriti & 24, Em. Fili str., GR-543 52, Thessaloniki,

**Facility Address** 

Product

SIDAPHARM BSS Sterile Balanced Salt Solution

500 ml Ref No: 76000 / 250 ml Ref No: 76001

GMDN: 37207

Classification

IIb, according to Rule 8, Annex IX of Council Directive

93/42/EEC

**Guidelines Applicable:** 

MEDDEV 2.4/1 Rev. 9 - June 2010 Rule 5

This Declaration of Conformity is issued under the sole responsibility of the manufacturer. SIDAPHARM hereby declares that the aforementioned product complies with all the essential requirements of Council Directive 93/42/EEC, concerning Medical Devices, as amended by 2007/47/EC.

The compliance of this applicable quality assurance system has been certified by the "Health Technology Certification", which is a Notified Body, according to Council Directive 93/42/EEC, with identification number 2803.

The present is issued, according to EC Certificate No.: 1828C04210505, whose original expiry date was 03/06/2023, but it has been granted an extension, according to Regulation (EU) 2023/607, till 31/12/2028.

The present replaces any previous declaration has been issued for these products.

For and on behalf of: SIDAPHARM P.C.

## SIDAPHARM P.C.

Medical Disposables Head Offices: 21, Stageiriti & 24, Em. Fili str. The station iki GR, 54852, Greece Branch: 6, Lacktod Str., Pylly GR-55535, Greece Tel: +30 2310 906660 - Fax: +30 2310 989846 VAT Reg. No.: EL997296038 Registration Number: 144520204000

Diana Mochintra General Manager

Date: 04/04/2024



## **ANNEX I – Applied Standards**

Standard Number	Title
EN ISO 13485:2016	Medical devices - Quality management systems - Requirements
EN 130 13403.2010	for regulatory purposes
EN ISO 14971:2019	Medical devices - Application of risk management to medical devices
EN ISO 11607-1:2020	Packaging for terminally sterilized medical devices -Part 1: Requirements for materials, sterile barrier systems and packaging systems
EN ISO 11607-2:2020	Packaging for terminally sterilized medical devices - Part 2: Validation requirements for forming, sealing and assembly processes
EN ISO 10993-1:2020	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process
EN 556-1:2001/AC:2006	Sterilization of medical devices – Requirements for medical devices to be designated "STERILE" - Part-1 Requirements for terminally sterilized medical devices
EN 62366:2015 / AC:2020	Medical devices - Application of usability engineering to medical devices
EN ISO 15223-1:2021	Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements
EN ISO 20417:2021	Information supplied by the manufacturer of medical devices
ISO 24971:2020	Medical devices - Guidance on the application of ISO 14971
EN ISO 16671:2015	Ophthalmic implants - Irrigating solutions for ophthalmic surgery
EN ISO 11138-1:2017	Sterilization of health care products - Biological indicators - Part 1: General requirements
EN ISO 11138-3:2017	Sterilization of health care products - Biological indicators - Part 3: Biological indicators for moist heat sterilization processes
ASTM D4169-16	Standard Practice for Performance Testing of Shipping Containers and Systems
EN ISO 14644-1:2015	Cleanrooms and associated controlled environments — Part 1: Classification of air cleanliness by particle concentration
EN ISO 14644-2:2015	Cleanrooms and associated controlled environments — Part 2: Monitoring to provide evidence of cleanroom performance related to air cleanliness by particle concentration
EN ISO 14644-3:2019	Clean rooms and associated controlled environments - Test methods
EN ISO 14644-4:2022	Clean rooms and associated control environments - Design, construction and start - up
EN ISO 17665-1:2006	Sterilization of health care products- Moist heat - Requirements for the development validation & routine control of sterilization process for medical device
ISO 14698-1:2003	Clean rooms and associated controlled environment - Biocontamination control - General principles and methods
ISO 14698 -2:2003	Cleanrooms and associated controlled environments - Biocontamination control Part 2: Evaluation and interpretation of biocontamination data