

## **EC Declaration of Conformity**

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Version	I
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## 1. EC Declaration of Conformity

Manufacturer : Rossmax Swiss GmbH

Address Widnauerstrasse 1, CH-9435 Heerbrugg, Switzerland

**Notified Body** SGS Belgium NV

Address SGS House Noorderlaan 87 2030 Antwerp Belgium

EU Identification No. : 1639

Certificate No. TW19/20056

Representative in Europe

: CMC Medical Devices& Drugs S.L. Address

C/ Horacio Lengo No 18, CP 29006, Málaga, Spain

Product type : Nasal irrigator

Type Designation : NW1

**Conformity Assessment** : EU Council Directive 93/42/EEC amended by 2007/47/EC Annex

II (excluding Section 4) and Regulation (EU) 2023/607 amendment of Regulations (EU) 2017/745 as regards the transitional provisions for

certain medical devices

Classification : Class IIa (According to EU Council Directive 93/42/EEC

amended by 2007/47/EC, Annex IX, Rule 11)

Lot No. YY M 001

▶ Lot sequential number

Manufacturing month.

Manufacturing year.

The above-mentioned devices are in full compliance with the relevant provisions of EU Council Directive 93/42/EEC amended by 2007/47/EC, Regulation (EU) 2023/607 amendment of Regulations (EU) 2017/745, Annex I-Essential Requirements and applied harmonized standards, national standards or other normative documents.

EN ISO 15223-1: 2016, EN 1041: 2008, EN ISO 10993-1:2009/ AC2010, EN ISO 10993-5:2009, EN ISO 10993-10:2013, EN ISO

13485:2016, EN ISO 14971:2019, EN 60601-1:2006/ A1:2013, EN 60601-1-2:2015, EN 60601-1-6:2010, EN62366:2008, EN60601-1-11

2010

Rossmax Swiss GmbH is exclusively responsible for the declaration of conformity.

This declaration is limited by the issuing of a revised declaration of conformity after change of the product and/or the expiration date (31.12.2028) of the certificate of 93/42/EEC.

Signature: Yolanda Lin, Management Representative

Date: Jun. 13, 2023