

DECLARATION OF CONFORMITY

Manufacturer's Name: Maxtec, LLC

Address: 2305 South 1070 West

Salt Lake City, Utah 84119

USA

SRN: US-MF-000003961

European Representative: EMERGO EUROPE

Westervoortsedijk 60, 6827 AT Arnhem The Netherlands

European Representative SRN: NL-AR-000000116

Basic UDI DI: 081777002MAXVENTURIFV

Product Name(s): MaxVenturi Family (MaxVenturi, Venturi Muffler)

Part Number/ Identification
R211P03-019
R211P03-020
R211P03-022
R211P03-024
R211P03-025
R211P03-026
R211P03-027
R211P03-033
R211P03-034
R211P03-036
R211P03-124
R211P38
R211P38-001

Intended Purpose: The MaxVenturi Series is intended for use in high-flow oxygen therapy

applications where the ability to deliver diluted oxygen to the breathing gas



pathway is required. Oxygen being delivered from this device is for adult patients. It is a restricted medical device intended for use by qualified, trained personnel, under the direction of a physician, in professional health care settings, i.e., hospital, sub-acute institutions, home care, and intra-

hospital transport.

The MaxVenturi is not a life supporting device.

GMDN: 44225 – Oxygen/air breathing gas mixer, hospital

Risk Class & Classification Class IIa Clause 3.2 Rule 10 of Annex IX of MDD

Criteria: Class IIa Clause 6.4 Rule 12 of Annex VIII MDR 2017/745

This EU declaration of conformity is issued under the sole responsibility of the manufacturer. We herewith declare that the above mentioned products meet the provisions of the following EC Council Regulations, Common Specifications and Standards. All supporting documents are retained under the premises of the manufacturer and the European Representative.

Regulations: General Application Directives: Medical Device Directive, Council

Directive 93/42/EEC of 14 June 1993 per Annex II Excluding Section 4 as

amended by 2007/47/EC of September 2007

RoHs (EU) 2015/863

Notified Body: TÜV SÜD Product Service

RIDLERSTRASSE 65, D-80339 MUNICH, Germany

Number 0123

Conformity Assessment: Full Quality Assurance per Annex II MDD 93/42/EEC Excluding Section

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EC Certificate No.: G1 045041 00 25 Rev. 00

Extension Confirmation Letter: CL 045041 0026

Date CE mark was affixed: 10 October 2007

This Declaration is considered valid from the date of the signature below until December 31, 2028.

Issued by Signature: _____ Date: 23 January 2025

Name: Anvitha Anand Rao

Position: Regulatory and Quality Product Engineering Specialist



Approved by Signature Date.	Approved by Signature: _		Date:	Jan 27,	2025
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Name: Rebecca Hudson

Position: VP, Regulatory Affairs and Quality Assurance

This Declaration supersedes any preceding Declaration of Conformity for the MaxVenturi Family.

Applied Common Specification and Standards:

The referenced list of Common Specifications and harmonized standards for which documented evidence of compliance can be provided includes:

EN ISO 13485:2016 (ISO 13485:2016) EN ISO 14971:2019 (ISO 14971:2019) EN ISO 20417:2021 ISO 15223-1:2021