

DECLARATION OF CONFORMITY

Manufacturer's Name:

Maxtec

Address:

2305 South 1070 West

Salt Lake City, Utah 84119

USA

European Representative:

QNET BV

Kantstraat19

NL-5076 NP Haaren The Netherlands

Product:

MAXO2+ Series Oxygen Analyzers

Model(s):

MAXO2+ A, MAXO2+ AE, and Handi+ (Including Pole Mount Bracket

Accessories R217P23, R205P86, R100P10, R206P76, and R206P75)

Classification & GMDN:

IIa Analyzer, Gas, Oxygen 35219

Classification criteria:

Clause 3.2 Rule 10 of Annex IX of MDD

We herewith declare that the above mentioned products meet the provisions of the following EC Council Directives and Standards. All supporting documents are retained under the premises of the manufacturer and the European Representative.

Directives:

General application directives: Medical Device Directive, COUNCIL DIRECTIVE

93/42/EEC of 14 June 1993 per Annex II as amended by 2007/47/EC of 5 September 2007

Notified Body:

TÜV SÜD Product Service

RIDLERSTRASSE 65, D-80339 MUNICH, Germany

Number 0123

EC Certificate No.:

G1 16 10 45041 020

Date CE mark was affixed:

21 June 2004, - MAXO2+/ 03 December 1998 - Handi+

This declaration is considered valid from April 29, 2019 to December 18, 2021.

Signature:

Date: 29 April 2019

Name:

Tammy Lavery

Position:

Director of Regulatory and Quality



Applied Standards

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

EN ISO 14971:2012 IEC 60601-1:2005 (EN 60601-1:2006) IEC 60601-1-2:2004 (EN 60601-1-2:2004) EN 62366:2008 EN 1041:2008 ISO 15223-1:2016