

DECLARATION OF CONFORMITY

Manufacturer's Name:

Maxtec

Address:

6526 South Cottonwood Street Salt Lake City, Utah 84107

USA

European Representative:

ONET BV

Hommerterweg 286 6436 AM Amstenrade The Netherlands

Product:

Oxygen Sensor

Model(s):

MAX-250 Series, MAX-2, MAX-8, MAX-25 & MAX-50

Classification & GMDN:

IIa

Sensor, Oxygen - 13538

Classification criteria:

Clause 1.2 Rule 2 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

Notified Body:

TÜV SÜD Product Service -123

RIDLERSTRASSE 65, D-80339 MUNICH, Germany

EC Certificate No.:

G1 14 06 45041 014

Date CE mark was affixed:

15 June 2000

This declaration is considered valid from December 19, 2011 to December, 18, 2016.

1 among Lavery Date: 9-2-2014

Name:

Tammy Lavery

Position:

Director of Regulatory & Quality/Maxtec Management Representative

TF0010 SECTION 13 REV K RF192 REV E



Applied Standards

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

ISO 13485:2012 ISO 7767:1967 EN ISO 14971:2012 EN 62366:2008 EN 1041:2008 ISO 15223-1:2012