

EC Declaration of Conformity

Manufacturer: DeVilbiss Healthcare LLC

100 DeVilbiss Drive

Somerset, PA 15501, USA

EC Authorized Representative: DeVilbiss Healthcare GmbH

Kamenzerstraße 3, 68309 Mannheim, Germany

1. Oxygen Concentrators (UMDNS 12-873)

Catalogue nos.: 525KS, 525KS-LT, 525KS-UK, 525PS
Description: DeVilbiss® 5-Liter Oxygen Concentrator

Classification (MDD Annex IX): IIa (Rule 11)

Conformity Assessment Procedure: MDD 93/42/EEC, Annex II excluding Section 4

2. Accessories:

Product Description (Catalogue no.):

Caster, 4 pkg.-DFT 501DZ-603 Flow Meter Pkg., Low Output 515LF-607 Sieve Bed SSP 525D-619 EXT Life Intake Bacteria Filter-DFT MC44D-605 Compressor Filter (Sintered Bronze) 525DD-626 Compressor SSP 525K-625 Compressor SSP 525PS-625 Cabinet Air Filter 303DZ-605 Final Bacteria Filter PV5LD-651 Oxygen outlet connector (plastic, 1/pack) CN100 Transfill Tubing PF1100TUB Transfill Caddy 525DD-650

Applied standards: All applicable harmonized standards as adopted under the EC Directive 93/42/EEC and published in the Official Journal of the European Communities (*See attached listing*).

This Declaration of Conformity is based on the EC Directive 93/42/EEC, Annex II.3 and supported by the TÜV NORD CERT GmbH (0044) certificate, with reference to articles 1 and 3 of the directive 93/42/EEC.

Notified Body: TÜV NORD CERT GmbH

Langemarckstrasse 20, 45141 Essen, Germany

Identification No.: 0044

EC Certificate No.: 44 232 117803 **Start of EC Marking:** 2007-10-01

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We, DeVilbiss Healthcare LLC., hereby declare under our sole responsibility that the above-mentioned products comply with the essential requirements and provisions of Council Directive 93/42/EEC.

Validity of this Declaration:

2019-08-07 - 2024-05-26

Somerset, PA,

Oct 26, 2022

Place Date

Roberto Munoz
Name and Position

Director, Regulatory Affairs & Audits

Applied Standards:

AAMI / ANSI ES60601-1:2005/(R) 2012 Ed 3.1 and A1:2012, C1:2009/(R) 2012 and A2:2010/(R) 2012 (Consolidated Text) Medical Electrical Equipment—Part 1: General Requirements for Basic Safety and Essential Performance (IEC 60601-1:2005, Mod). (General I (QS/RM))

AAMI / ANSI / IEC 60601–1–2:2014, Ed. 4.0, Medical Electrical Equipment—Part 1-2: General Requirements for Basic Safety And Essential Performance – Collateral Standard; Electromagnetic Disturbances - Requirements and Tests (associated with IEC 60601-1 Ed. 3.0)

IEC 60601-1-8 Ed. 2.1 b.2012 Medical electrical equipment - Part 1-8: General requirements for basic safety and essential performance - Collateral Standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems (associated with IEC 60601-1 3rd Edition)

IEC 60601-1-11:2015 Medical electrical equipment - Part 1-11: General requirements for basic safety and essential performance - Collateral Standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment (associated with IEC 60601-1 3rd Edition, referenced by ISO 80601-2-69:2014)

AAMI / ANSI / IEC 62304:2006, Medical Device Software – Software Life Cycle Process (Software/Informatics)

ISO 80601-2-69:2014 Medial electrical equipment – Part 2-69: Particular requirements for basic safety and essential performance of oxygen concentrator equipment

ISTA Procedure 3APackaged-Products for Parcel Delivery System Shipments 70kg (150 lb) or Less (standard)

AMMI / ANSI / ISO 10993-1:2009, Biological Evaluation Of Medical Devices – Part 1: Evaluation And Testing Within A Risk Management Process (Biocompatibility)

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