

EC Declaration of Conformity

DeVilbiss Healthcare LLC

100 DeVilbiss Drive
Somerset, PA 15501, USAEC Authorized Representative:DeVilbiss Healthcare GmbH
Kamenzerstraße 3, 68309
Mannheim, Germany1. Oxygen Delivery Units, controlled (UMDNS 18-076):

1. Oxygen Delivery Units, controlled (UMDNS 18-076):
Catalogue nos.:535/

Classification (MDD Annex IX):IIa (Rule 11)Conformity Assessment Procedure:MDD 93/42/EEC, Annex II excluding Section 4

2. Accessories:

Manufacturer:

Product Description (Catalogue no.):

ML6 Cylinder w/CF Regulator, Europe	5351-ML6-CF
C Cylinder w/CF Regulator, Europe	5351-C-CF
D Cylinder w/CF Regulator, Europe	535I-D-CF
E Cylinder w/CF Regulator, Europe	5351-E-CF
Cabinet Air Filter, 6 pk.	535D-605
Bag, C Cylinder	EX3000D-651
Bag, D Cylinder	EX3000D-652
Bag, M-6 Mini Cylinder	EX3000D-653
Bag, ML6 Cylinder	EX3000D-654
Cylinder Nipple Dust Cap, 5 pk.	PD1000A-627
Covers, Top & Bottom replacement	PD1000A-601
Gauge Kit	PD1000A-620
Gauge Kit	PD1000G-620
Seal, Regulator, 10 pk.	PD1000G-618
PD1000A w/C Cylinder, Europe	PD1000A-I-C
PD1000A w/D Cylinder, Europe	PD1000A-I-D
PD1000A w/E Cylinder, Europe	PD1000A-I-E
PD1000A w/ML6 Cylinder, Europe	PD1000A-I-ML6

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:	2002-06-19



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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.

Somerset, PA, <u>30 July 2024</u>

Place

Date

Zita A. Yurko, RAC Sr. Director Regulatory Affairs Name and Position

Applied Standards:

5 series	
C 60601-1:2005+A1:2012 Medial electrical equipment — Part 1 General requirements for basic safety	
C 60601-1-2:2014, Medical electrical equipment — Part 1-2: General requirements for basic safety and essentic rformance — Collateral Standard: Electromagnetic compatibility – Requirements and tests (FDA Recognition Imber 19-8)	1
C 60601-1-6:2010 + AMD 1:2013 Ed. 3.1 Medical electrical equipment - Part 1-6: General requirements for basi fety and essential performance - Collateral standard: Usability (associated with IEC 60601-1 Ed. 3.0) (FDA cognition Number 5-89)	С
C 60601-1-9:2007 + A1:2013 Ed. 1.1 Medical electrical equipment - Part 1-9: General requirements for basic saj d essential performance - Collateral standard: Requirements for environmentally conscious design (associated C 60601-1 Ed. 3.0)	-
C 60601-1-11:2015 (Ed 2.0), Medical electrical equipment — Part 1-11: General requirements for basic safety a sential performance – Collateral Standard: Requirements for medical electrical equipment and medical electric stems used in the home healthcare environment (FDA Recognition Number19-14)	
C 62366:2007 Ed. 1.0 + AMD 1:2014 – Medical devices - Application of usability engineering to medical devices DA Recognition number 5-87)	;
C 62304:2006 + AMD1:2015: Ed. 1.1 - Medical device software - Software life cycle processes (FDA Recognition mber 13-79)	
EN ISO 14971:2012 Medical devices - Application of risk management to medical devices (FDA Recognition mber 5-40 is to ISO 14971:2007	
A Procedure 3A: Packaged-Products for Parcel Delivery System Shipments 70kg (150 lb) or less (standard).	
TM G63-15 Standard Guide for Evaluating Nonmetallic Materials for Oxygen Service	
TM G94-05 Standard Guide for Evaluating Metals for Oxygen Service	
EN ISO 15001:2011 Anaesthetic and respiratory equipment. Compatibility with oxygen	