

# **EC Declaration of Conformity**

Manufacturer: DeVilbiss Healthcare LLC

100 DeVilbiss Drive

Somerset, PA 15501, USA

DeVilbiss Healthcare GmbH **EC Authorized Representative:** 

> Kamenzerstraße 3, 68309 Mannheim, Germany

1. Suction Units (UMDNS 13-846):

7314D-AP, 7314D-D, 7314D-D-EXF, 7314D-LA, 7314D-NE, 7314D-U, Catalogue nos.:

7314P-AP, 7314P-D, 7314P-D-EXF, 7314P-LA, 7314P-NE, 7314P-NE-R,

7314P-U, 7314P-UR

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

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

### 2. Accessories:

Product Description (Catalogue no.):

Collection Container Kit (internal filter cartridge, splash guard, 800 ml container, 4 ½" and 6" tubing package)  800 ml Disposable Container w/ internal filter cartridge, splash guard, 4 ½" tubing, 48 each  Filter Cartridge assy, 1 pk. (for 800 ml disposable container use) 7305D-634 Filter Cartridge, 12 pk. (for 800 ml disposable container use) 7305D-635 Collection Container Kit (1200 ml reusable container, external bacteria filter, elbow, 4 ½" tubing) 1200 ml Reusable Container (external bacteria filter, elbow, 4 ½" tubing) 6 pk. External Bacteria Filter (non-sterile), 12 pk. (for reusable container use) 7305D-608 Battery, w/ 2 pcs. 7314 Foam, replacement assy. 7314P-614
tubing, 48 each  Filter Cartridge assy, 1 pk. (for 800 ml disposable container use)  Filter Cartridge, 12 pk. (for 800 ml disposable container use)  Collection Container Kit (1200 ml reusable container, external bacteria filter, elbow, 4 %" tubing)  1200 ml Reusable Container (external bacteria filter, elbow, 4 %" tubing) 6 pk.  External Bacteria Filter (non-sterile), 12 pk. (for reusable container use)  7305D-634  7305D-635  7314D-603
Filter Cartridge, 12 pk. (for 800 ml disposable container use)  Collection Container Kit (1200 ml reusable container, external bacteria filter, elbow, 4 %" tubing)  1200 ml Reusable Container (external bacteria filter, elbow, 4 %" tubing) 6 pk.  External Bacteria Filter (non-sterile), 12 pk. (for reusable container use)  7305D-635  7314D-603
Collection Container Kit (1200 ml reusable container, external bacteria filter, elbow, 4 %" tubing)  1200 ml Reusable Container (external bacteria filter, elbow, 4 %" tubing) 6 pk.  External Bacteria Filter (non-sterile), 12 pk. (for reusable container use)  7314D-603  7314D-604  7305D-608
elbow, 4 %" tubing)  1200 ml Reusable Container (external bacteria filter, elbow, 4 %" tubing) 6 pk.  External Bacteria Filter (non-sterile), 12 pk. (for reusable container use)  7314D-604  7305D-608
External Bacteria Filter (non-sterile), 12 pk. (for reusable container use) 7305D-608
Battery, w/ 2 pcs. 7314 Foam, replacement assy. 7314P-614
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Carry Case 7314D-606
AC to DC adapter/charger 7314P-613
DC Power Cord, 1 each 7304D-619
Power Cord, USA DV51D-606
Power Cord, Continental Europe DV51D-607
Power Cord, UK DV51D-608
Power Cord, Australia DV51D-609
Power Cord, Brazil DV51D-612
Power Cord, Japan DV51D-613
Power Cord, China DV51D-614
Power Cord, Argentina 180-0006-011

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.

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## **EC Declaration of Conformity**

Notified Body: TÜV NORD CERT GmbH

Langemarckstrasse 20, 45141 Essen, Germany

Identification No.: 0044

**EC Certificate No.:** 44 232 117803 Start of EC Marking: 15-08-2012

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, MAI 7, 2023 Kabello L. M.

Roberto Munoz Director, Regulatory Affairs and Audit

Place, Date Nume and Positi

### **Applied Standards:**

#### 7314 series

EN ISO 10079-1:2015 Ed.3 Medical Suction Equipment-Part 1:Electrically Powered Suction Equipment IEC 60601-1:2005+A1:2012 Medial electrical equipment — Part 1 General requirements for basic safety (FDA Recognition Number1-115)

IFC 60601-1-2:2014, Medical electrical equipment — Part 1-2: General requirements for basic safety and essential performance — Collateral Standard: Electromagnetic compatibility – Requirements and tests (FDA Recognition Number 19 8)

IEC 60601-1-6:2010 + AMD 1:2013 Ed. 3.1 Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability (associated with IEC 60601-1 Ed. 3.0) (FDA Recognition Number 5-89)

IEC 60601-1-11:2015 (Ed 2.0), 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 (FDA Recognition Number19-14)

IEC 62366:2007 Ed. 1.0 + AMD 1:2014 — Medical devices - Application of usability engineering to medical devices (FDA Recognition number 5-87)

BS EN ISO 14971:2012 Medical devices - Application of risk management to medical devices (FDA Recognition number 5-40 is to ISO 14971:2007)

IEC 60529 Issued 2001/02/01 Ed:2.1, Classification of Degrees of Protection Provided by Enclosures
IEC 60068-2-6 Issued:2007/12/01 Ed:7.0 Environmental Testing-Part 2-6:Tests-Test Fc: Vibration (sinusoidal)
IEC 60068-2-27 Issued 2008/02/01 Ed:4.0 Environmental Testing-Part 2-27: Tests-Test Ea and guidance: Shock
IEC 60068-2-34 Issued 1973/01/01 Ed.1 Basic Environmental Testing Procedures Part 2: Tests Test Fd: Random Vibration Wide Band-General Requirements

ISTA 3A Packaged Product Testing: Dynamic Vibration, Drop Testing, Thermal Testing