

# EU DECLARATION OF CONFORMITY



Doc Number 2101660

Revision 05

This Declaration of Conformity is issued under the sole responsibility of the manufacturer. The device(s)/accessories covered by this Declaration are in conformity with all regulations or directives below.

## 1. Object of the declaration:

<b>Product Name:</b>	<i>SimplyGo Mini Accessories</i>	
<b>Product Type:</b>	<i>Portable Oxygen Concentrator Accessories</i>	
<b>Intended Purpose:</b>	<p><b>GMDN 34158 Battery, Secondary</b>                      A device (battery or cell) used as a source of electrical energy that is designed to be electrically recharged. The size, shape, and chemical composition of the battery should be specified in keeping with the requirements of the appropriate IEC standard.</p> <p><b>GMDN 17115 Battery Charger</b>                      A device designed to supply an electrical charge to rechargeable batteries, restoring the battery to an appropriate working condition. This device is typically connected to the building's electrical power supply and can be used to either charge the batteries by themselves (removed from the device) or whilst they are still inside the parent device (in situ), e.g. a defibrillator or ophthalmoscope. This device usually has current and voltage controls to meet the charge needs of different types of batteries.</p>	
<b>Product Part Number(s) and Descriptions:</b>	1119949	SimplyGo Mini Battery Charger, UK
	1119950	SimplyGo Mini Battery Charger, EU
	1116830	SimplyGo Mini Battery Charger, NA
	1116816	SimplyGo Mini Standard Battery Kit
	1116817	SimplyGo Mini Extended Battery Kit
	1129319	SimplyGo Mini, Extended Battery Kit, Saudi Arabia
	1129307	SimplyGo Mini Standard Battery Kit, Saudi Arabia
	FR1113604	SimplyGo Mini, Standard Battery, FR
	FR1113605	SimplyGo Mini, Extended Battery, FR
<b>Product Options/Accessories Part Number(s) and Descriptions:</b>	Accessories to the SimplyGo device  Note: SimplyGo DoC: REG 2101229	
<b>Basic UDI-DI:</b>	1119949	606959403277
	1119950	606959403291
	1116830	606959403239
	1116816	606959403468
	1116817	606959403475
	1129319	606959407473
	1129307	606959407480
	FR1113604	606959058453

**CONFIDENTIAL**

This document was created using the template information listed below:

<b>Governing Document:</b> GSP 7.9-064, WI 7.9-808	<b>Document Number:</b> FRM 4450	<b>Version:</b> 10	Page 1 of 4
---	----------------------------------	--------------------	-------------

# EU DECLARATION OF CONFORMITY



Doc Number 2101660  
Revision 05

	FR1113605 606959058460	
<b>Control Indicator:</b>	Initial Issue Date	Part Number
	January 18, 2017	1129319, 1129307
	October 6, 2015	1119949, 1119950, 1116830, 1116816, 1116817
	See Date Below	FR1113604, FR1113605
<b>EMDN / CND code and description And/or Global Medical Device Nomenclature code (GMDN) and Description:</b>	17115 - External device battery charger (1119949, 1119950, 1116830) 34158 - Secondary battery (1116816, 1116817, 1129319, 1129307. FR1113604, FR1113605)	

The object of the declaration described above is in conformity with the following regulations:

<b>EU Directive</b>	<b>Council Directive 93/42/EEC of 14 June 1993 concerning medical devices amended up to and inclusive of Council Directive 2007/47/EC (MDD)</b>
<b>Risk Classification</b>	Class IIa, Rule 2
<b>Conformity Assessment Route</b>	Annex II excluding (4)
<b>Notified Body Name, Address, and ID</b>	TÜV SÜD Product Service GmbH Ridlerstrasse 65 80339 München, Germany 0123
<b>Certificate(s) Issued</b>	EC certificate: G1 015581 068
<b>Standards</b>	The products listed on this Declaration of Conformity have been assessed and/or tested in a typical configuration as described in the Manufacturer's accompanying documentation in accordance with the product standards listed below.  Refer to Attachment A.

**CONFIDENTIAL**

This document was created using the template information listed below:

<b>Governing Document:</b> QSP 7.9-064, WI 7.9-808	<b>Document Number:</b> FRM 4450	<b>Version:</b> 10	Page 2 of 4
---	----------------------------------	--------------------	-------------

# EU DECLARATION OF CONFORMITY



Doc Number 2101660

Revision 05

## 2. Additional information:

<b>Manufacturer</b>	Respironics, Inc. 1001 Murry Ridge Lane, Murrysville, PA 15668, USA
<b>EU Authorized Representative:</b>	Respironics Deutschland GmbH & Co. KG Gewerbestrasse 17 82211 Herrsching, Germany Tel: +49 8152 93060
<b>ISO Quality Certificates Issued:</b>	The Manufacturer is certified by TÜV SÜD Product Service GmbH to the following:  EN ISO 13485 Certificate: Q5 015581 0607 MDSAP ISO 13485 Certificate: QS6 17 10 15581 058

Signature (signed for and on behalf of  
*Respironics, Inc.*)

Date of Issue: September 24, 2020

Printed Name: Daria Brown

Place of Issue: Monroeville, PA, USA

Title: Sr. Manager, Regulatory Affairs

**CONFIDENTIAL**

This document was created using the template information listed below:

Governing Document:  
QSP 7.9-064, WI 7.9-808

Document Number: FRM 4450

Version: 10

Page 3 of 4

# EU DECLARATION OF CONFORMITY



Doc Number 2101660  
Revision 05

## 3. Attachment A Standards and/or Common Specifications

Standard	Standard Title
<b>Quality System</b>	
EN ISO 13485:2016	Medical devices – Quality management systems – Requirements for regulatory purposes
<b>General Safety Standard</b>	
EN 60601-1:2006/A1:2013	Medical electrical equipment -- Part 1: General requirements for basic safety and essential performance
<b>Collateral Safety Standards</b>	
EN 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
<b>Particular Safety Standards</b>	
<b>Biocompatibility</b>	
EN ISO 10993-1:2018	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process
<b>Other Standards</b>	
<b>Accompany Documents and Labeling</b>	
EN 1041:2008	Information supplied by the manufacturer of medical devices
EN ISO 15223-1:2017	Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements
<b>Risk Management</b>	
EN ISO 14971:2019	Medical devices – Application of risk management to medical devices
<b>Common Specifications</b>	

### CONFIDENTIAL

This document was created using the template information listed below:

Governing Document:  
QSP 7.9-064, WI 7.9-808

Document Number: FRM 4450

Version: 10

Page 4 of 4