

# EU DECLARATION OF CONFORMITY



Doc Number REG 2101485  
Revision 04

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:

|  |  |   |                    |                    |         |   |         |                                   |
|--|--|---|--------------------|--------------------|---------|---|---------|-----------------------------------|
| Product Name:  | SimplyGo Accessories   |   |                    |                    |         |   |         |                                   |
| Product Type:  | Portable Oxygen Concentrator Accessories   |   |                    |                    |         |   |         |                                   |
| Intended Purpose:  | <p>The SimplyGo Batteries are accessories for the SimplyGo and share its intended purpose. The Respironics SimplyGo Portable Oxygen Concentrator is for prescription use by patients requiring high concentrations of oxygen on a supplemental basis. It is small, portable, and is capable of continuous use in the home, institutional, and travel/mobile environments.</p> <p>The Philips Respironics SimplyGo External Battery Module can be used to power the SimplyGo device with a SimplyGo rechargeable Lithium-ion battery. When a SimplyGo battery is inserted into the module and the module is connected to the SimplyGo, the module will extend the battery duration of the device. When the module is connected, the SimplyGo will initially use the external battery power. Once the external battery is fully depleted, the SimplyGo will automatically switch over to its internal battery. The battery used with the External Battery Module can be recharged using your SimplyGo device or the Philips Respironics Smart Charger.</p> |   |                    |                    |         |   |         |                                   |
| Product Part Number(s) and Descriptions:                     | <p>Part Number(s) listed in this section comply with all regulation and directive indicated in DoC unless otherwise noted.</p> <table><tr><td><u>Part Number</u></td><td><u>Description</u></td></tr><tr><td>1082662</td><td>SimplyGo, Battery (Extra / Replacement)</td></tr><tr><td>1109643</td><td>SimplyGo, External Battery Module</td></tr></table>  |   | <u>Part Number</u> | <u>Description</u> | 1082662 | SimplyGo, Battery (Extra / Replacement) | 1109643 | SimplyGo, External Battery Module |
| <u>Part Number</u>   | <u>Description</u>   |   |                    |                    |         |   |         |                                   |
| 1082662  | SimplyGo, Battery (Extra / Replacement)  |   |                    |                    |         |   |         |                                   |
| 1109643  | SimplyGo, External Battery Module  |   |                    |                    |         |   |         |                                   |
| Product Options/Accessories Part Number(s) and Descriptions: | None   |   |                    |                    |         |   |         |                                   |
| Basic UDI-DI:  | 0606959BM153LW   |   |                    |                    |         |   |         |                                   |
| Control Indicator:   | <u>Initial Issue Date:</u><br>August 8, 2017<br><br>May 21, 2014   | <u>Part Number:</u><br>1082662<br><br>1109643 |                    |                    |         |   |         |                                   |

### 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: 11

Page 1 of 5

# EU DECLARATION OF CONFORMITY



Doc Number REG 2101485

Revision 04

|   |  |
|---|--|
| <b>EMDN / CND code and Description</b>                                | CND:<br>Z12159004 Oxygen Concentrators |
| <b>And</b>  |  |
| <b>Global Medical Device Nomenclature code (GMDN) and Description</b> | GMDN:<br>34158 - Secondary battery     |

The object of the Declaration (product part numbers and if applicable the component part numbers) that are described above as included in this DoC is / are in conformity with the following regulations / directives. If optional accessories that are used in combination with the product (medical device), but are CE marked on their own are referenced in this DoC, they are excluded further on as these optional accessories have their own DoC.

|  |   |
|--|---|
| <b>EU Regulation</b>                       | <b>Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices (EU MDR)</b>   |
| <b>Risk Classification</b>                 | Class I based on Annex VIII and Rule 13   |
| <b>Conformity Assessment Route</b>         | N/A   |
| <b>Notified Body Name, Address, and ID</b> | TÜV SÜD Product Service GmbH<br>Ridlerstrasse 65<br>80339 München, Germany<br><br>NB ID – Not Applicable  |
| <b>Certificate(s) Issued</b>               | N/A   |
| <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.<br><br>Refer to Attachment A |
| <b>Common Specifications</b>               | None  |

## CONFIDENTIAL

This document was created using the template information listed below:

|   |                                  |                    |                    |
|---|----------------------------------|--------------------|--------------------|
| <b>Governing Document:</b><br>QSP 7.9-064, WI 7.9-808 | <b>Document Number:</b> FRM 4450 | <b>Version:</b> 11 | <b>Page</b> 2 of 5 |
|---|----------------------------------|--------------------|--------------------|



# EU DECLARATION OF CONFORMITY



Doc Number REG 2101485

Revision 04

|                            |   |
|----------------------------|---|
| <b>EU Directive</b>        | <b>Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restrictions of the use of certain hazardous substances in electrical and electronic equipment, amended up to and inclusive of Directive (EU) 2017/1202 (RoHS)</b>      |
| <b>Risk Classification</b> | Category 8, medical device, according Annex I   |
| <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.<br><br>Refer to Attachment A |

## 2. Mandatory information:

|   |  |
|---|--|
| <b>Manufacturer</b>                       | Respironics, Inc.<br>1001 Murry Ridge Lane,<br>Murrysville, PA 15668, USA<br><br>Legal Manufacturer SRN: US-MF-000002301   |
| <b>EU Authorized Representative (AR):</b> | Philips Medical Systems Nederland B.V.<br>Veenpluis 6 5684PC Best, The Netherlands<br><br>Single Registration Number (SRN): NL-AR-000001422  |
| <b>ISO Quality Certificates Issued:</b>   | The Manufacturer is certified by TÜV SÜD Product Service GmbH to the following:<br><br>EN ISO 13485 as evidenced by Q5 015581 0609<br><br>ISO 13485 MDSAP Certificate Number:<br>QS6 015581 0610<br><br><i>Copies of the Quality System certificates are available upon request.</i> |

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

Printed Name: Nicole Beale

Date of Issue: 21 June 2021

Place of Issue: Monroeville, PA, USA

Title: Sr. Manager, Regulatory Affairs

| CONFIDENTIAL   |                                  |                    |                    |
|--|----------------------------------|--------------------|--------------------|
| This document was created using the template information listed below: |                                  |                    |                    |
| <b>Governing Document:</b><br>QSP 7.9-064, WI 7.9-808                  | <b>Document Number:</b> FRM 4450 | <b>Version:</b> 11 | <b>Page</b> 3 of 5 |

# EU DECLARATION OF CONFORMITY



Doc Number REG 2101485

Revision 04

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

| Standard                           | Standard Title   | Product Applicability |
|------------------------------------|--|-----------------------|
| <b>Quality System</b>              |  |                       |
| <b>EN ISO 13485:2016</b>           | Medical devices – Quality management systems – Requirements for regulatory purposes  | All Products          |
| <b>General Safety Standard</b>     |  |                       |
| <b>EN 60601-1:2006/A1:2013</b>     | Medical electrical equipment -- Part 1: General requirements for basic safety and essential performance  | All Products          |
| <b>Collateral Safety Standards</b> |  |                       |
| <b>EN 60601-1-2:2015</b>           | Medical electrical equipment – Part 1-2: General requirements for basic safety and essential performance – Collateral standard: Electromagnetic compatibility. Requirements and tests  | All Products          |
| <b>EN 60601-1-6:2010/A1:2015</b>   | Medical electrical equipment – Part 1-6: General requirements for safety and essential performance – Collateral standard: Usability  | All Products          |
| <b>EN 60601-1-8:2007/A1:2013</b>   | Medical electrical equipment – Part 1-8: General requirements for basic safety and essential performance – Collateral Standard: General requirements, tests and guidance for alarm systems used in medical electrical equipment and medical electrical systems | All Products          |
| <b>EN 60601-1-11:2015</b>          | 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          | All Products          |
| <b>Particular Safety Standards</b> |  |                       |
| <b>Oxygen Concentrators</b>        |  |                       |
| <b>EN ISO 80601-2-69:2014</b>      | Medical Electrical Equipment — Part 2-69: Particular requirements for basic safety and essential performance of oxygen concentrator equipment  | All Products          |

### CONFIDENTIAL

This document was created using the template information listed below:

|   |                                  |                    |                    |
|---|----------------------------------|--------------------|--------------------|
| <b>Governing Document:</b><br>QSP 7.9-064, WI 7.9-808 | <b>Document Number:</b> FRM 4450 | <b>Version:</b> 11 | <b>Page 4 of 5</b> |
|---|----------------------------------|--------------------|--------------------|



# EU DECLARATION OF CONFORMITY



Doc Number REG 2101485  
Revision 04

|   |  |                  |
|---|--|------------------|
| <b>Oxygen Conserving Devices</b>        |  |                  |
| <b>ISO 80601-2-67:2014</b>              | Medical electrical equipment — Part 2-67: Particular requirements for basic safety and essential performance of oxygen conserving equipment  | All Products     |
| <b>Batteries</b>                        |  |                  |
| <b>IEC 62133:2012</b>                   | Secondary cells and batteries containing alkaline or other non-acid - electrolytes — Safety requirements for portable sealed secondary cells, and for batteries made from them, for use in portable applications | SimplyGo Battery |
| <b>Other Standards</b>                  |  |                  |
| <b>Accompany Documents and Labeling</b> |  |                  |
| <b>EN 1041:2008/A1:2013</b>             | Information supplied by the manufacturer of medical devices  | All Products     |
| <b>EN ISO 15223-1:2017</b>              | Medical Devices – Symbols to be used with medical device labels, labelling and information to be supplied. Part 1: General requirements  | All Products     |
| <b>Risk Management</b>                  |  |                  |
| <b>EN ISO 14971:2019</b>                | Medical devices – Application of risk management to medical devices  | All Products     |
| <b>Usability</b>                        |  |                  |
| <b>IEC 62366-1:2015</b>                 | Medical devices – Part 1: Application of usability engineering to medical devices  | All Products     |
| <b>RoHS</b>                             |  |                  |
| <b>EN IEC 63000:2018</b>                | Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances   | All Products     |

## CONFIDENTIAL

This document was created using the template information listed below:

|   |                                  |                    |                    |
|---|----------------------------------|--------------------|--------------------|
| <b>Governing Document:</b><br>QSP 7.9-064, WI 7.9-808 | <b>Document Number:</b> FRM 4450 | <b>Version:</b> 11 | <b>Page</b> 5 of 5 |
|---|----------------------------------|--------------------|--------------------|