

Doc Number REG 2101735 Revision 08

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:	DreamWisp Nasal Mask with Over the Nose Cushion			
Product Type:	Nasal Mask			
Intended Purpose:	This nasal mask is intended to provide an interface for application of Continuous Positive Airway Pressure (CPAP) or bi-level therapy to patients. The mask is intended for single patient use in the home or multi-patient use in the hospital/institutional environment. The mask is to be used on patients (>66 lbs/30kg) for whom CPAP or bi-level therapy has been prescribed.			
Product Part Number(s) and Descriptions:	1137916 FitPack, DreamWisp Nsl, Med Con w/Hgr GBL 1137918 FitPack, DreamWisp Nsl, Med Con w/o Hgr GBL 1137922 P, DreamWisp Nsl, Med Con w/o Hgr GBL			
	1137923 S, DreamWisp Nsl, Med Con w/o Hgr GBL 1137924 M, DreamWisp Nsl, Med Con w/o Hgr GBL 1137925 L, DreamWisp Nsl, Med Con w/o Hgr GBL 1137926 XL, DreamWisp Nsl, Med Con w/o Hgr GBL			
	1137932 P, DreamWisp Nsl, Med Con w/Hgr, GBL 1137933 S, DreamWisp Nsl, Med Con w/Hgr, GBL 1137934 M, DreamWisp Nsl, Med Con w/Hgr, GBL 1137935 L, DreamWisp Nsl, Med Con w/Hgr, GBL 1137936 XL, DreamWisp Nsl, Med Con w/Hgr, GBL			
	1137953 LABPACK, DREAMWISP, ALL CON, S/M/L CUSH W/HGR 1137954 P/XL LABPK, DREAMWISP LAB PACK, S/L CON COMBO LAB PACK, DREAMWISP			
	1137921 FitPack, DreamWisp Nsl, Med Con w/Hgr INTL 1137942 P, DreamWisp Nsl, Med Con w/Hgr, INTL 1137943 S, DreamWisp Nsl, Med Con w/Hgr, INTL 1137944 M, DreamWisp Nsl, Med Con w/Hgr, INTL 1137945 L, DreamWisp Nsl, Med Con w/Hgr, INTL 1137946 XL, DREAMWISP NSL, MED CON W/HGR, INT			
	1137942AP P, DreamWisp Nsl, Med Con, W/HGR, AP 1137943AP S, DreamWisp Nsl, Med Con, W/HGR, AP 1137944AP M, DreamWisp Nsl, Med Con, W/HGR, AP			

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	1137945AP 1137946AP 1137942CE 1137943CE 1137945CE 1137946CE 1137943RC 1137943RC 1137945RC 1137946RC 1137946RC	P, Dream S, Dream M, Dream L, Dream XL, Dream S, Dream M, Dream L, Dream XL, Dream	Wisp Nsl, Med Con, W/HGR,AP mWisp Nsl, Med Con, W/HGR, CE Wisp Nsl, Med Con, W/HGR, CE mWisp Nsl, Med Con, W/HGR, CE wisp Nsl, Med Con, W/HGR, RC
Product Options/Accessories Part Number(s) and Descriptions:			neadgear and chinstrap accessories
Basic UDI-DI:	N/A		
Control Indicator:	Initial Issue 16 January, 27 March, 20 2 April, 2019	2019 019 9	Part Number: 1137916, 1137918, 1137922, 1137923, 1137924, 1137925, 1137926, 1137932, 1137933, 1137934, 1137935, 1137936 1137953, 1137954 1137943, 1137944, 1137942, 1137946 1137942AP, 1137943AP,
	02 June, 202	20	1137944AP, 1137945AP, 1137946AP, 1137942CE, 1137943CE, 1137944CE, 1137945CE, 1137946CE, 1137942RC, 1137943RC, 1137944RC, 1137945RC, 1137946RC
Global Medical Device Nomenclature code (GMDN) and Description	•		sal Mask Reusable

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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.

EU Directive	Council Directive 93/42/EEC of 14 June 1993 concerning medical devices amended up to and inclusive of Council Directive 2007/47/EC (MDD)
Risk Classification	Class IIa based on Annex IX and Rule 2
Conformity Assessment Route	Annex II excluding (4)
Notified Body Name, Address, and ID	TÜV SÜD Product Service GmbH Ridlerstrasse 65 80339 München, Germany Notified Body Number: 0123
Certificate(s) Issued	TÜV SÜD EC Certificate Number: G1 015581 0611
Standards	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.

#### 2. Mandatory information:

Manufacturer	Respironics, Inc.		
	1001 Murry Ridge Lane		
	Murrysville, PA 15668		
	USA		
EU Authorized	Respironics Deutschland GmbH & Co. KG		
Representative (AR):	Gewerbestrasse 17		
	82211 Herrsching, Germany		
	Tel: +49 8152 93060		
ISO Quality	The Manufacturer is certified by TÜV SÜD to the following:		
Certificates Issued:	TÜV SÜD EN ISO 13485:2016 Certificate Number: Q5		
	015581 0609		
	TÜV SÜD MDSAP Certificate Number: QS6 112601 0001		

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

Date of Issue: 27 August 2021

Printed Name: Jennifer Richardson

Place of Issue: Murrysville, PA

Title: Senior Manager, Regulatory Affairs

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3. Attachment A Standards and/or Common Specifications

Standard	Standard Title			
Quality System				
EN ISO 13485:2016	Medical devices – Quality management systems – Requirements for regulatory purposes			
Particular Safety Standards				
Patient Interface				
EN ISO 17510-2:2009	Sleep apnoea breathing therapy - Part 2: Masks and application accessories			
EN ISO 17510:2020	Medical devices - Sleep apnoea breathing therapy - Masks and application accessories			
Biocompatibility				
EN ISO 10993-1:2020	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process			
EN ISO 10993-3:2014	Biological evaluation of medical devices - Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity			
EN ISO 10993-5:2009	Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity			
EN ISO 10993-6:2016	Biological evaluation of medical devices - Part 6: Tests for local effects after implantation			
EN ISO 10993-10:2013	Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization			
EN ISO 10993-11:2018	Biological evaluation of medical devices - Part 11: Tests for systemic toxicity			
EN ISO 10993-17:2009	Biological evaluation of medical devices - Part 17: Establishment of allowable limits for leachable substances			
EN ISO 10993-18:2009	Biological evaluation of medical devices - Part 18: Chemical characterization of materials			
ISO 18562-1:2017	Biocompatibility evaluation of breathing gas pathways in healthcare applications - Part 1: Evaluation and testing within a risk management process			
ISO 18562-2:2017	Biocompatibility evaluation of breathing gas pathways in healthcare applications - Part 2: Tests for emissions of particulate matter			
ISO 18562-3:2017	Biocompatibility evaluation of breathing gas pathways in healthcare applications - Part 3: Tests for emissions of volatile organic compounds (VOCs)			
ISO 18562-4:2017	Biocompatibility evaluation of breathing gas pathways in healthcare applications - Part 4: Tests for leachables in condensate			
Other Standards				
Accompany Documents and Labeling				

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Standard	Standard Title		
Quality System			
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		
EN 1041:2008/A1:2013	Information supplied by the manufacturer of medical devices		
Risk Management			
ISO 14971:2019	Medical devices - Application of risk management to medical devices		
Usability			
IEC 62366-1: 2015	Medical devices – Part 1: Application of usability engineering to medical devices		
Cleaning and Disinfection			
EN ISO 17664:2017	Processing of health care products - Information to be provided by the medical device manufacturer for the processing of medical devices		
Tubing and Connections			
EN ISO 5356-1:2015	Anaesthetic and respiratory equipment - Conical connectors - Part 1: Cones and sockets		

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