



## Declaration of Conformity

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Manufacturer: *ResMed Ltd*  
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Bella Vista  
NSW 2153  
Australia

European Representative: *ResMed (UK) Ltd*  
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Abingdon  
Oxfordshire OX14 4RY  
United Kingdom

Notified Body: *SGS United Kingdom Ltd*  
Weston-Super-Mare  
Unit 202b, Worle Parkway  
Weston-Super-Mare, BS22 0WA  
United Kingdom



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**Product:** **Swift LT**

The Swift LT channels airflow noninvasively to a patient from a positive airway pressure device such as a continuous positive airway pressure (CPAP) or bilevel system.

The Swift LT is:

- to be used by adult patients (> 66 lb/30 kg) for whom positive airway pressure has been prescribed.
- intended for single-patient re-use in the home environment and multipatient re-use in the hospital/institutional environment.

Standards Applied: ISO 17510-2:2007;  
ISO 10993-1:2003; ISO 5356-1:2004 (sections 5.1 and 5.2);  
ISO 594-1:1986 (sections 3 and 4.1); ISO 17664:2004.

Classification: Ila

Conformity

Assessment Route: Annex II (excluding Section 4), 93/42/EEC

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We herewith declare that the above mentioned products meet the transposition into national law of the provisions of Council Directive 93/42/EEC including the MDD amendment (2007/47/EC), for medical devices. All supporting documentation is retained at the premises of the manufacturer.

Sydney, *9<sup>th</sup> May 2008.*

Dr. Lionel King  
Vice President Global Quality Assurance and Regulatory Affairs  
ResMed Ltd

*Director, Regulatory Affairs. ResMed Ltd.*

**EC093**