

Document title

Declaration of Conformity

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We, Atos Medical AB, hereby declare that the below mentioned devices comply with Swedish regulation LVFS 2003:11, transposing Annex II, with the exemption of section 4 (for class IIa and class IIb devices) and Annex VII (for class I devices) of the European Medical Devices Directive 93/42/EEC and clause 3.5 of Schedule 3 to the Therapeutic Goods (Medical Devices) Regulations 2002.

The Trach-HME Products

REF	Name	Class	GMDN code
7704	TrachPhone (50 pcs)	lla	58705
7705	MEDIFLUX HCH F6 (Medival)	lla	58705
7707	TrachPhone (30 pcs)	lla	58705
7723	TrachPhone (5 pcs)	lla	58705

Each kind of medical device to which the system has been applied complies with the applicable provisions of the Essential Principles and Essential Requirements, the classification rules, applicable standards and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

Valid Certificates are controlled and filed by QA/RA Department. For standards applied see Applicable Standards & Essential Requirements for each product.

Notified Body: Intertek Semko AB, Sweden. Identification no. 0413

EC-certificate no. 41310296-04

Competent Authority: Medical Products Agency, Sweden

Atos Medical AB Kraftgatan 8

SE-242 35 Hörby, Sweden Tel: +46 (0) 415 198 00

E-mail: info@atosmedical.com

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Approval Task Verdict: Approve	SEHRBHNU Ulrika Svensson, Regulatory Affairs Professional (ulrika.svensson-atosmedical@coloplast.com) Issuer 22-Nov-2024 09:12:21 GMT+0000
Approval Task Verdict: Approve	SEHRBPNH Håkan Persson, Quality Manager (hakan.persson-atosmedical@coloplast.com) Quality 22-Nov-2024 09:44:20 GMT+0000
Approval Task Verdict: Approve	MARRIC Martin Richardson, VP, International Manufacturing, Sweden, Germany & China (martin.richardson-atosmedical@coloplast.com) Management 06-Dec-2024 07:27:07 GMT+0000

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