



# CERTIFICATE



This is to certify that the company

## Medikro Oy

Pioneerinkatu 3 70800 Kuopio Finland

with the organizational units/sites as listed in the annex

has implemented and maintains a Quality Management System.

Scope of certification:

Design and development, manufacturing, distribution and servicing of pulmonary function measurement devices and their accessories.

- AUS (a), CND, USA (a,b,c,d)

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of the following standard:

# ISO 13485 : 2016

including applicable country-specific regulatory requirements, as indicated below the scope (full references of abbreviations are listed in the annex)

Certificate registration no.	542490 MDSAP16
Certificate unique ID	170781819
Effective date	2023-05-08
Expiry date	2025-05-30
Frankfurt am Main	2023-05-08

#### DQS Medizinprodukte GmbH

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Sigrid Uhlemann Managing Director



Marc Goedecke

Marc Goedecke Product Manager



August-Schanz-Straße 21, 60433 Frankfurt am Main, Tel. +49 (0) 69 95427-300, <u>info-med@dqs.de</u> **DQS Medizinprodukte GmbH is recognized under the Medical Devices Single Audit Program.** Visit <u>https://www.dqs.de/en/customer-database/</u> to validate this certificate. **The validity of this certificate can only be verified by the QR-code**.





#### Annex to certificate Certificate registration No.: 542490 MDSAP16 Certificate unique ID: 170781819 Effective date: 2023-05-08

## Medikro Oy

Pioneerinkatu 3 70800 Kuopio Finland

Audited site

REPs FEI No.: site scope and country-specific requirements

542490 Medikro Oy Pioneerinkatu 3 70800 Kuopio Finland

Design and development, manufacturing, distribution and servicing of pulmonary function measurement devices and their accessories. - AUS (a), CND, USA (a,b,c,d) REPs FEI No.: F003496





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Full references of country-specific requirements of MDSAP participating Regulatory Authorities

Abbreviation	Jurisdiction	Reference
AUS	Australia	<ul> <li>(a) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 1 – Full Quality Assurance Procedure</li> <li>(b) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 4 – Production Quality Assurance Procedure</li> </ul>
BRA	Brazil	RDC ANVISA n. 665/2022 RDC ANVISA n. 551/2021 RDC ANVISA n. 67/2009
CND	Canada	Medical Device Regulations SOR/98-282, Part 1
JPN	Japan	MHLW Ministerial Ordinance No. 169 (2004) as amended by MHLW Ordinance No. 128 (2014), Articles 4 to 68 Japan PMD Act (as applicable)
USA	United States	<ul> <li>(a) 21 CFR Part 803</li> <li>(b) 21 CFR Part 806</li> <li>(c) 21 CFR Part 807</li> <li>(d) 21 CFR Part 820</li> <li>(e) 21 CFR Part 821</li> </ul>

This annex is only valid in connection with the above-mentioned certificate.