CERTIFICATE OF REGISTRATION



Safey Medical Devices Private Limited

Plot No.PAP-S-47 &48, Phase - II, MIDC Sawardari, Tal- Khed, Chakan Pune, Maharashtra 410501 INDIA

Facility ID: F005403

UL Medical Regulatory Services of UL LLC®(UL) issues this certificate to the Firm named above, after auditing the Firm's quality management system and finding it in conformance per the defined scope with respect to:

ISO 13485:2016

with additional regulatory requirements listed on final page of this certificate.

Design, development and manufacture of spirometers, peak flow meters and mouthpiece for areas of anesthesiology.

MDSAP MEDICAL DEVICE SINGLE AUDIT PROGRAM Authorized by

Paul Hilgeman

Director & Global Industry Leader, Medical
CMIT – Medical Regulatory

Camp By Prince (i)

Check Certificate Status:

here

File Number A28913 Cycle Start Date June 11, 2021
Certificate Number 3313.230611 Effective Date June 11, 2023
Initial Issue Date June 11, 2021 Expiry Date June 10, 2024

This quality system registration is included in UL's Directory of Registered Firms and applies to the provision of goods and/or services as specified in the scope of registration from the address(es) shown above. By issuance of this certificate the firm represents that it will maintain its registration in accordance with the applicable requirements. This certificate is not transferable and remains the property of UL Medical and Regulatory Services of UL LLC. Certificates may be verified by visiting the Online Certifications Directory on UL.com.



UL Medical and Regulatory Services UL, LLC is an MDSAP Recognized Auditing Organization

UL LLC 333 Pfingsten Road Northbrook, IL 60062-2096 USA

CERTIFICATE OF REGISTRATION



Safey Medical Devices Private Limited

Plot No.PAP-S-47 &48, Phase - II, MIDC Sawardari, Tal- Khed, Chakan Pune, Maharashtra 410501 INDIA

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Additional Regulatory Requirements

Australia:

- Therapeutic Goods (Medical Devices) Regulations, 2002, Schedule 3 Part 1 (excluding Part 1.6) – Full Quality Assurance Procedure

Brazil:

- RDC ANVISA n. 665/2022
- RDC ANVISA n. 551/2021
- RDC ANVISA n. 67/2009

Canada:

- Medical Devices Regulations - Part 1- SOR 98/282

United States:

- 21 CFR 820
- 21 CFR 803
- 21 CFR 806
- 21 CFR 807 Subparts A to D
- 21 CFR 821 (where applicable)

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