



Institut luxembourgeois de la normalisation
de l'accréditation, de la sécurité et qualité
des produits et services

ILNAS-EN ISO 18778:2022

Respiratory equipment - Particular requirements for basic safety and essential performance of infant cardiorespiratory monitors (ISO

Matériel respiratoire - Exigences
particulières relatives à la sécurité de
base et aux performances essentielles
des moniteurs cardiorespiratoires pour

Medizinische elektrische Geräte -
Besondere Festlegungen für die
Sicherheit einschließlich der
wesentlichen Leistungsmerkmale von

10/2022



National Foreword

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**Respiratory equipment - Particular requirements for basic
safety and essential performance of infant
cardiorespiratory monitors (ISO 18778:2022)**

Matériel respiratoire - Exigences particulières relatives
à la sécurité de base et aux performances essentielles
des moniteurs cardiorespiratoires pour nourrissons
(ISO 18778:2022)

Medizinische elektrische Geräte - Besondere
Festlegungen für die Sicherheit einschließlich der
wesentlichen Leistungsmerkmale von
kardiorespiratorischen Überwachungsgeräten für
Kleinkinder (ISO 18778:2022)

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European foreword

This document (EN ISO 18778:2022) has been prepared by Technical Committee ISO/TC 121 "Anaesthetic and respiratory equipment" in collaboration with Technical Committee CEN/TC 215 "Respiratory and anaesthetic equipment" the secretariat of which is held by BSI.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by April 2023, and conflicting national standards shall be withdrawn at the latest by April 2023.

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The text of ISO 18778:2022 has been approved by CEN as EN ISO 18778:2022 without any modification.

Respiratory equipment — Particular requirements for basic safety and essential performance of infant cardiorespiratory monitors

Matériel respiratoire — Exigences particulières relatives à la sécurité de base et aux performances essentielles des moniteurs cardiorespiratoires pour nourrissons

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