



Institut luxembourgeois de la normalisation
de l'accréditation, de la sécurité et qualité
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ILNAS-EN ISO 80601-2-13:2022

Medical electrical equipment - Part 2-13: Particular requirements for basic safety and essential performance of an anaesthetic workstation (ISO

Medizinische elektrische Geräte - Teil
2-13: Besondere Festlegungen für die
Sicherheit einschließlich der
wesentlichen Leistungsmerkmale von

Appareils électromédicaux - Partie 2-13:
Exigences particulières de sécurité de
base et de performances essentielles
pour les postes de travail d'anesthésie

06/2022



National Foreword

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Medizinische elektrische Geräte - Teil 2-13: Besondere Festlegungen für die Sicherheit einschließlich der wesentlichen Leistungsmerkmale für Anästhesie-Arbeitsplätzen (ISO 80601-2-13:2022)

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European foreword

This document (EN ISO 80601-2-13:2022) 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.

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