



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

ILNAS-EN ISO 23500-5:2024

Preparation and quality management of fluids for haemodialysis and related therapies - Part 5: Quality of dialysis fluid for haemodialysis and related

Herstellung und Qualitätsmanagement
von Flüssigkeiten für die Hämodialyse
und verwandte Therapien - Teil 5:
Qualität von Flüssigkeiten für die

Préparation et management de la qualité
des liquides d'hémodialyse et de
thérapies annexes - Partie 5: Qualité des
liquides de dialyse pour hémodialyse et

04/2024



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**Preparation and quality management of fluids for
haemodialysis and related therapies - Part 5: Quality of
dialysis fluid for haemodialysis and related therapies (ISO
23500-5:2024)**

Préparation et management de la qualité des liquides
d'hémodialyse et de thérapies annexes - Partie 5:
Qualité des liquides de dialyse pour hémodialyse et
thérapies apparentées (ISO 23500-5:2024)

Herstellung und Qualitätsmanagement von
Flüssigkeiten für die Hämodialyse und verwandte
Therapien - Teil 5: Qualität von Flüssigkeiten für die
Hämodialyse und verwandte Therapien (ISO 23500-
5:2024)

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European foreword

This document (EN ISO 23500-5:2024) has been prepared by Technical Committee ISO/TC 150 "Implants for surgery" in collaboration with Technical Committee CEN/TC 205 "Non-active medical devices" the secretariat of which is held by DIN.

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of an identical text or by endorsement, at the latest by October 2024, and conflicting national standards shall be withdrawn at the latest by October 2024.

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The text of ISO 23500-5:2024 has been approved by CEN as EN ISO 23500-5:2024 without any modification.



**Preparation and quality
management of fluids for
haemodialysis and related
therapies —**

**Part 5:
Quality of dialysis fluid for
haemodialysis and related therapies**

*Préparation et management de la qualité des liquides
d'hémodialyse et de thérapies annexes —*

*Partie 5: Qualité des liquides de dialyse pour hémodialyse et
thérapies apparentées*



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