



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

ILNAS-EN ISO 23500-4:2024

Preparation and quality management of fluids for haemodialysis and related therapies - Part 4: Concentrates for haemodialysis and related therapies

Herstellung und Qualitätsmanagement
von Flüssigkeiten für die Hämodialyse
und verwandte Therapien - Teil 4:
Konzentrate für die Hämodialyse und

Préparation et management de la qualité
des liquides d'hémodialyse et de
thérapies annexes - Partie 4: Concentrés
pour hémodialyse et thérapies

04/2024



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**Preparation and quality management of fluids for
haemodialysis and related therapies - Part 4: Concentrates
for haemodialysis and related therapies (ISO 23500-
4:2024)**

Préparation et management de la qualité des liquides
d'hémodialyse et de thérapies annexes - Partie 4:
Concentrés pour hémodialyse et thérapies apparentées
(ISO 23500-4:2024)

Herstellung und Qualitätsmanagement von
Flüssigkeiten für die Hämodialyse und verwandte
Therapien - Teil 4: Konzentrate für die Hämodialyse
und verwandte Therapien (ISO 23500-4:2024)

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European foreword

This document (EN ISO 23500-4:2024) 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.

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ILNAS-EN ISO 23500-4:2024

International Standard

ISO 23500-4

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

Part 4: Concentrates for haemodialysis and related therapies

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

Partie 4: Concentrés pour hémodialyse et thérapies apparentées

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ISO copyright office
CP 401 • Ch. de Blandonnet 8
CH-1214 Vernier, Geneva
Phone: +41 22 749 01 11
Email: copyright@iso.org
Website: www.iso.org

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