



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

## ILNAS-EN ISO 8536-4:2020

### **Infusion equipment for medical use - Part 4: Infusion sets for single use, gravity feed (ISO 8536-4:2019)**

Infusionsgeräte zur medizinischen  
Verwendung - Teil 4: Infusionsgeräte für  
Schwerkraftinfusionen zur einmaligen  
Verwendung (ISO 8536-4:2019)

Matériel de perfusion à usage médical -  
Partie 4: Appareils de perfusion non  
réutilisables, à alimentation par gravité  
(ISO 8536-4:2019)

01/2020



## National Foreword

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ILNAS-EN ISO 8536-4:2020

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Infusionsgeräte zur medizinischen Verwendung - Teil  
4: Infusionsgeräte für Schwerkraftinfusionen zur  
einmaligen Verwendung (ISO 8536-4:2019)

This European Standard was approved by CEN on 17 October 2019.

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**CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels**

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## European foreword

This document (EN ISO 8536-4:2020) has been prepared by Technical Committee ISO/TC 76 "Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use" 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of an identical text or by endorsement, at the latest by July 2020, and conflicting national standards shall be withdrawn at the latest by July 2020.

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The text of ISO 8536-4:2019 has been approved by CEN as EN ISO 8536-4:2020 without any modification.

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Part 4:  
Infusion sets for single use, gravity feed**

*Matériel de perfusion à usage médical —*

*Partie 4: Appareils de perfusion non réutilisables, à alimentation  
par gravité*



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