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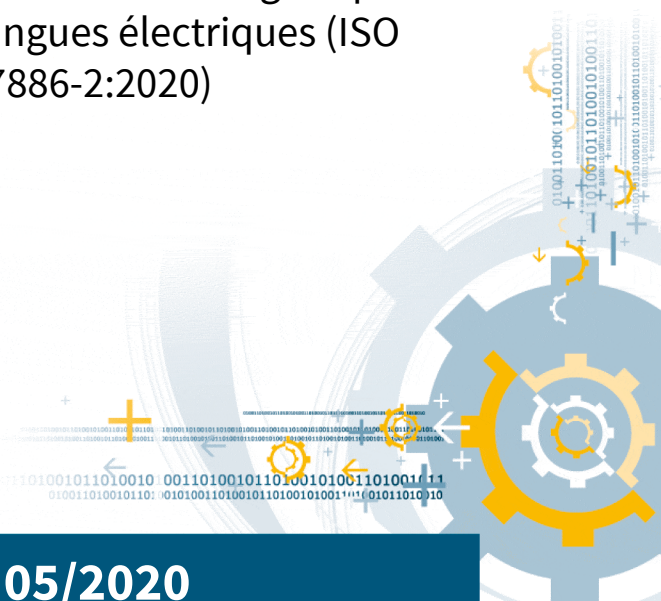
ILNAS-EN ISO 7886-2:2020

Sterile hypodermic syringes for single use - Part 2: Syringes for use with power-driven syringe pumps (ISO 7886-2:2020)

Sterile Einmalspritzen für medizinische
Zwecke - Teil 2: Spritzen zur Verwendung
mit Spritzenpumpen (ISO 7886-2:2020)

Seringues hypodermiques stériles, non
réutilisables - Partie 2: Seringues pour
pousse-seringues électriques (ISO
7886-2:2020)

05/2020



National Foreword

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EUROPEAN STANDARD **EN ISO 7886-2**

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English Version

**Sterile hypodermic syringes for single use - Part 2:
Syringes for use with power-driven syringe pumps (ISO
7886-2:2020)**

Seringues hypodermiques stériles, non réutilisables -
Partie 2: Seringues pour pousse-seringues électriques
(ISO 7886-2:2020)

Sterile Einmalspritzen für medizinische Zwecke - Teil
2: Spritzen zur Verwendung mit Spritzenpumpen (ISO
7886-2:2020)

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European foreword

This document (EN ISO 7886-2:2020) has been prepared by Technical Committee ISO/TC 84 "Devices for administration of medicinal products and catheters" 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of an identical text or by endorsement, at the latest by November 2020, and conflicting national standards shall be withdrawn at the latest by November 2020.

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Endorsement notice

The text of ISO 7886-2:2020 has been approved by CEN as EN ISO 7886-2:2020 without any modification.

Sterile hypodermic syringes for single use —

Part 2: Syringes for use with power-driven syringe pumps

*Seringues hypodermiques stériles, non réutilisables —
Partie 2: Seringues pour pousse-seringues électriques*



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