



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

ILNAS-EN ISO 20342-1:2022

Assistive products for tissue integrity when lying down - Part 1: General requirements (ISO 20342-1:2022)

Produits d'assistance pour l'intégrité des
tissus en position allongée - Partie 1:
Exigences générales (ISO 20342-1:2022)

Hilfsmittel für die Gewebeintegrität im
Liegen - Teil 1: Allgemeine Anforderungen
(ISO 20342-1:2022)

08/2022



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ILNAS-EN ISO 20342-1:2022

EUROPEAN STANDARD **EN ISO 20342-1**

NORME EUROPÉENNE

EUROPÄISCHE NORM

August 2022

ICS 11.180.01

Supersedes EN ISO 20342-1:2019

English Version

**Assistive products for tissue integrity when lying down -
Part 1: General requirements (ISO 20342-1:2022)**

Produits d'assistance pour l'intégrité des tissus en
position allongée - Partie 1: Exigences générales (ISO
20342-1:2022)

Hilfsmittel für die Gewebeintegrität im Liegen - Teil 1:
Allgemeine Anforderungen (ISO 20342-1:2022)

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European foreword

This document (EN ISO 20342-1:2022) has been prepared by Technical Committee ISO/TC 173 "Assistive products" in collaboration with Technical Committee CEN/TC 293 "Assistive products and accessibility" the secretariat of which is held by SIS.

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The text of ISO 20342-1:2022 has been approved by CEN as EN ISO 20342-1:2022 without any modification.

Assistive products for tissue integrity when lying down —

Part 1: General requirements

*Produits d'assistance pour l'intégrité des tissus en position
allongée —*

Partie 1: Exigences générales



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Published in Switzerland

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