



Technical Specification

ISO/TS 20721

Implants for surgery — Absorbable implants — General guidelines and requirements for assessment of absorbable metallic implants

*Implants chirurgicaux — Implants absorbables — Lignes
directrices et exigences générales pour l'évaluation des implants
métalliques absorbables*

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Foreword

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This document was prepared by Technical Committee ISO/TC 150, *Implants for surgery*.

This second edition cancels and replaces the first edition (ISO/TS 20721:2020), which has been technically revised.

The main change is as follows: references have been updated.

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