
**Implants for surgery — Components
for partial and total knee joint
prostheses —**

**Part 2:
Articulating surfaces made of metal,
ceramic and plastics materials**

AMENDMENT 2

*Implants chirurgicaux — Éléments de prothèses partielle et totale de
l'articulation du genou —*

*Partie 2: Surfaces articulaires constituées de matériaux métalliques,
céramiques et plastiques*

AMENDEMENT 2 38-846e-18c240fc5799/iso-7207-2-2011-amd-2-2020



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Published in Switzerland

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This document was prepared by Technical Committee ISO/TC 150, *Implants for surgery*, Subcommittee SC 4, *Bone and joint replacements*.

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