
**Implants for surgery — Wear of
total intervertebral spinal disc
prostheses —**

Part 1:
**Loading and displacement parameters
for wear testing and corresponding
environmental conditions for test**

AMENDMENT 1

*Implants chirurgicaux — Usure des prothèses totales de
remplacement des disques intervertébraux lombaires —*

*Partie 1: Paramètres de charge et de déplacement pour essais d'usure
et conditions environnementales correspondantes*

AMENDEMENT 1



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