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Cardiovascular implants and artificial organs — Hard-shell cardiectomy/venous reservoir systems (with/without filter) and soft venous reservoir bags

Implants cardiovasculaires et organes artificiels — Systèmes réservoirs de cardiectomie/veineux à paroi dure (avec/sans filtre) et sacs réservoirs veineux mous



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ISO copyright office
Ch. de Blandonnet 8 • CP 401
CH-1214 Vernier, Geneva, Switzerland
Tel. +41 22 749 01 11
Fax +41 22 749 09 47
copyright@iso.org
www.iso.org

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Foreword

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The committee responsible for this document is ISO/TC 150, *Implants for surgery*, Subcommittee SC 2, *Cardiovascular implants and extracorporeal systems*.

This third edition cancels and replaces the second edition (ISO 15674:2009), which has been technically revised.