
**Intravascular catheters — Sterile and
single-use catheters —**

**Part 6:
Subcutaneous implanted ports**

AMENDMENT 1

Cathéters intravasculaires — Cathéters stériles et non réutilisables —

Partie 6: Chambres à cathéter implantables

AMENDEMENT 1

ISO 10555-6:2015/Amd 1:2019

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