



International Standard

ISO 5361

Anaesthetic and respiratory equipment — Tracheal tubes and connectors

AMENDMENT 1: Reinstatement of third edition S1 dimensions

*Matériel d'anesthésie et de réanimation respiratoire — Sondes
trachéales et raccords*

*AMENDEMENT 1: Rétablissement des dimensions S1 de la
troisième édition*

**Fourth edition
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