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AMENDMENT 1
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**Flow-metering devices for connection
to terminal units of medical gas
pipeline systems**

AMENDMENT 1

*Dispositifs de mesure de débit pour raccordement aux prises murales
des systèmes de distribution de gaz médicaux*

AMENDEMENT 1

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ISO 15002:2008/Amd 1:2018

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This document was prepared by Technical Committee ISO/TC 121, *Anaesthetic and respiratory equipment*, Subcommittee SC 6, *Medical gas systems*.

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