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AMENDMENT 1
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**Anaesthetic and respiratory
equipment — Low-pressure hose
assemblies for use with medical gases**

AMENDMENT 1

*Matériel d'anesthésie et de réanimation respiratoire — Flexibles de
raccordement à basse pression pour utilisation avec les gaz médicaux*

AMENDEMENT 1

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This document was prepared by Technical Committee ISO/TC 121, *Anaesthetic and respiratory equipment*, Subcommittee SC 1, *Breathing attachments and anaesthetic machines*.

Introduction

This document has been prepared in response to the revision of ISO 7396-1, which applies in its third edition also to oxygen supply systems in which all sources of supply deliver oxygen 93. Following this decision, ISO/TC 121/SC 1 approved developing the 1st amendment to ISO 5359:2014 by replacing the term “oxygen-enriched air” by “oxygen 93” and by introducing the designation “O₂ 93” for the medical gas oxygen 93.

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