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Anaesthetic and respiratory equipment-_— General requirements for airway devices and related equipment

iTeh Standards (https://standards.iteh.ai)

Matériel d'anesthésie et de réanimation respiratoire — Exigences générales pour canules et équipement <u>connexe</u>

<u>ISO/FDIS 18190</u>

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Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO document should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT), see www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 121, *Anaesthetic and respiratory equipment*, Subcommittee SC 2, *Airway devices and related equipment*.

<u>ISO/FDIS 18190</u>

This second edition cancels and replaces the first edition (ISO 18190:2016), which has been technically 76365a5de0/iso-fdis-18190 revised.

The main changes are as follows:

- — Title altered from airways to airway devices.
- The introduction has been changed to clarify that this standard can be used in the absence of a device specific standard.
- ----Definitions for clinical evaluation and clinical investigation added.
- — Risk management process and *clinical evaluation* now mandated.
- A new requirement recommending that manufacturers consider the environmental impact of their device and its packaging during its lifetime has been added.
- — A requirement for the biological evaluation for devices with breathing gas pathways has been added
- Information provided by the manufacturer, including marking, now refers to ISO 20417 for the common requirements and only lists those requirements specific to airway devices and related equipment.

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- Devices that are safe, conditionally safe or unsafe to be used in an MR environment are now to be marked accordingly.
- The requirements for positioning of controls and protection against inadvertent adjustments have been deleted as they were deemed not applicable to airway devices.
- A new requirement for shelf life has been added.
- All requirements relating to sterility have been condensed into one clause.
- A new requirement has been added for cleanliness and disinfection and combined with sterility.
- A new requirement to disclose the transport and environmental conditions that the airway device can withstand has been added.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at <u>www.iso.org/members.html</u>www.iso.org/members.html.

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Introduction

This document provides the general requirements for basic safety and performance for _the design, materials, packaging, marking and labelling that are generally applicable to all *airway devices and* related equipment.

This document is intended to consolidate the general requirements that are common among the set of standards within the category of *airway devices and related equipment* and serve as a reference for these common requirements, allowing each device-specific standard to focus on the unique safety and essential requirements more concisely for that device.

This document should be used in conjunction with device-specific *airway devices and related equipment* standards.

The requirements in a device-specific standard take priority over any conflicting requirements in this document.

If there is no airway device-specific standard, then this document can be referenced for all the applicable requirements.

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NOTE The terms defined in Clause 3 are denoted throughout the document in *italic font*.

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Anaesthetic and respiratory equipment ----- General

requirements for airway devices and related equipment

1 Scope

This document specifies the general requirements common to airway devices and related equipment.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO-4135, Anaesthetic and respiratory equipment — Vocabulary

 $ISO_{-10993-1}$, Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process

ISO-11607-1, <u>Packaging for terminally sterilized medical devices — Part 1:</u> Requirements for materials, sterile barrier systems and packaging systems

ISO-11607-2, Packaging for terminally sterilized medical devices — Part 2: Validation requirements for forming, sealing and assembly processes — Amendment 1: Application of risk management

ISO-<u>11135</u>, Sterilization of health-care products — Ethylene oxide — Requirements for the development, validation and routine control of a sterilization process for medical devices for development, validation and routine control of a sterilization process for medical devices/Amendment <u>1</u>

ISO-11137-1, Sterilization of health care products — Radiation — Part 1: Requirements for the development, validation and routine control of a sterilization process for medical devices

ISO-14155:2020, Clinical investigation of medical devices for human subjects — Good clinical practice

ISO-14937:2009, Sterilization of health care products — General requirements for characterization of a sterilizing agent and the development, validation and routine control of a sterilization process for medical devices

ISO-14971, Medical devices — Application of risk management to medical devices

ISO-15001:2010, Anaesthetic and respiratory equipment — Compatibility with oxygen

ISO-15223-1:2021, Medical devices — Symbols to be used with information to be supplied by the manufacturer — <u>Part 1:</u> General requirements

ISO–17664-1, Processing of health care products — Information to be provided by the medical device manufacturer for the processing of medical devices — Part 1: Critical and semi-critical medical devices

ISO-17664-2, Processing of health care products — Information to be provided by the medical device manufacturer for the processing of medical devices — Part 2: Non-critical medical devices

ISO-17665, Sterilization of health care products — Moist heat — Requirements for the development, validation and routine control of a sterilization process for medical devices

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ISO-_18562-1, Biocompatibility evaluation of breathing gas pathways in healthcare applications — Part 1: Evaluation and testing within a risk management process

ISO-<u>1</u>8601, Packaging and the environment — General requirements for the use of ISO standards in the field of packaging and the environment

ISO-20417, Medical devices — informationInformation to be supplied by the manufacturer

ISO-20857, Sterilization of health care products — Dry heat — Requirements for the development, validation and routine control of a sterilization process for medical devices

ISO-22441, Sterilization of health care products — Low temperature vaporized hydrogen peroxide — Requirements for the development, validation and routine control of a sterilization process for medical devices

ISO-25424, Sterilization of health care products — Low temperature steam and formaldehyde — Requirements for development, validation and routine control of a sterilization process for medical devices — Amendment 1

ISO-80369-7, Small-bore connectors for liquids and gases in healthcare applications — Part 7: Connectors for intravascular or hypodermic applications

IEC-60601-1:2005, + AMD1:2012 + AMD2:2020, Medical electrical equipment — Part 1: General requirements for basic safety and essential performance

IEC-60601-1-2:2014+AMD1:2020, Medical electrical equipment — Part 1-2: General requirements for basic safety and essential performance — Collateral standard: Electromagnetic disturbances — Requirements and tests

IEC-60601-1-8:2006, + AMD1:2012 + AMD2:2020, Medical electrical equipment — Part 1-8: General requirements for basic safety and essential performance _— Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems

IEC-62570:2014, Standard practice for marking medical devices and other items for safety in the magnetic resonance environment

ASTM F640, Standard test methods for determining radiopacity for medical use

EN-556-1⁺, Sterilization of medical devices. Requirements for medical devices to be designated "__STERILE". Part 1. Requirements for terminally sterilized medical devices-

EN-15986, Symbol for use in the labelling of medical devices — Requirements for labelling of medical devices containing phthalates

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 4135, ISO 14971, ISO 20417 and the following apply.

ISO and IEC maintain terminology databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <u>https://www.iso.org/obp</u>
- IEC Electropedia: available at <u>https://www.electropedia.org/</u>

Field Code Changed

⁴ Under development at enquiry stage at time of publication of this document.

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