

Czech metrology institute Notified Body No. 1383

Český metrologický institut Oznámený subjekt č. 1383

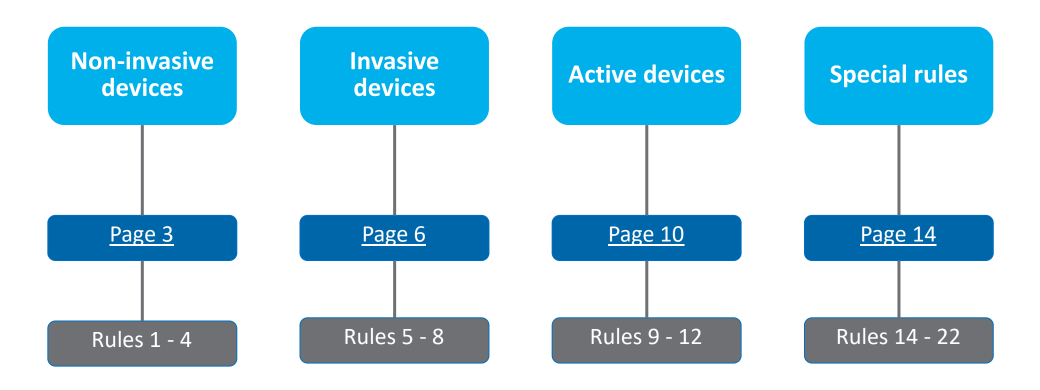


Methodological guide

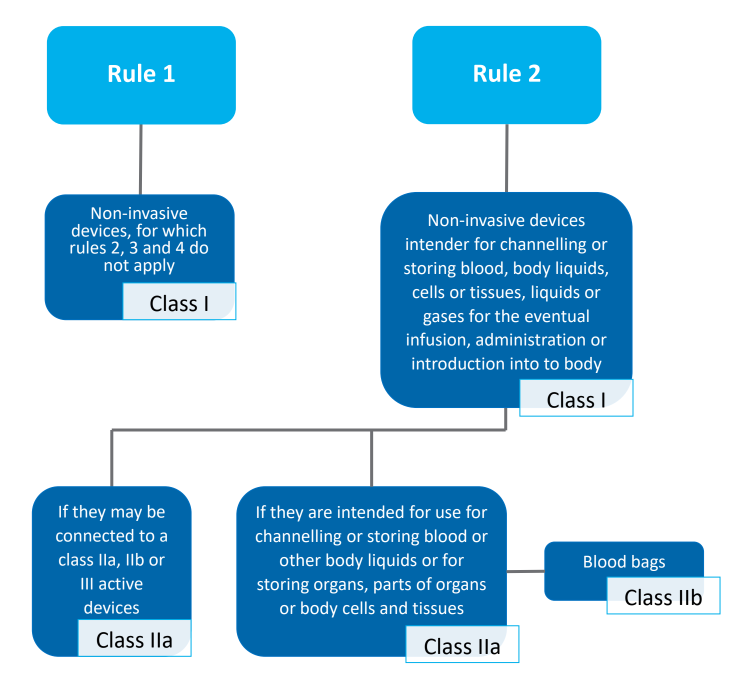
MEDICAL DEVICES CLASSIFICATION RULES

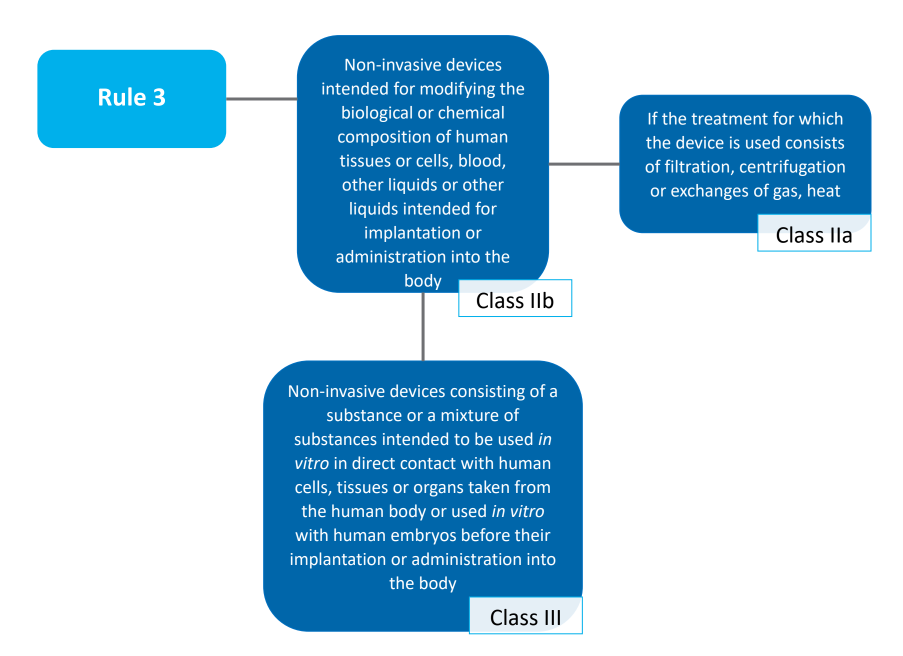
This methodological guide is only an aid for orientation in classification rules for medical devices. To determine the class of your medical device we recommend using Annex VIII of the MDR Regulation.

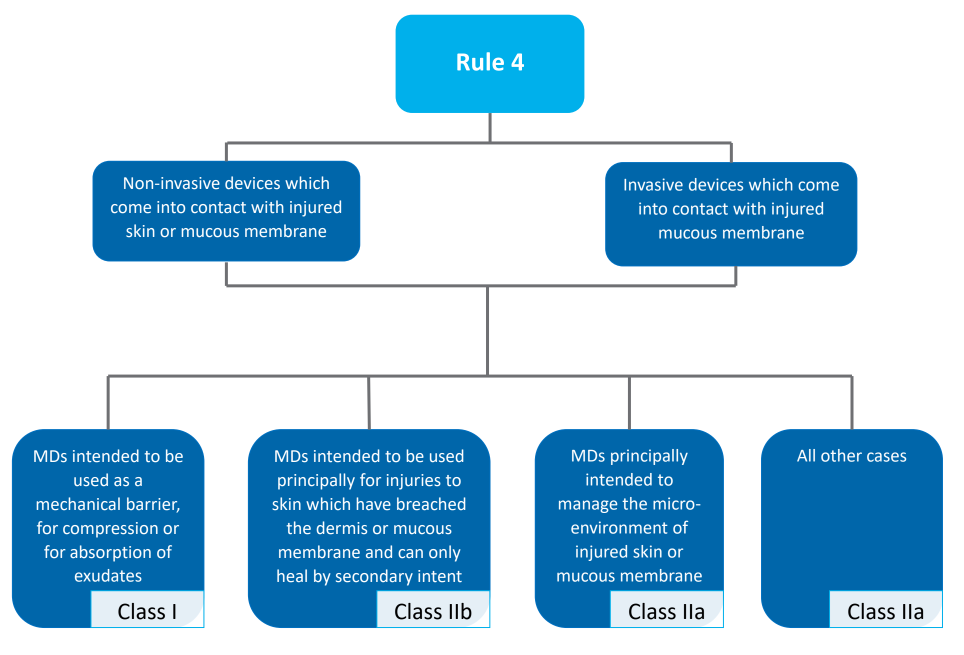


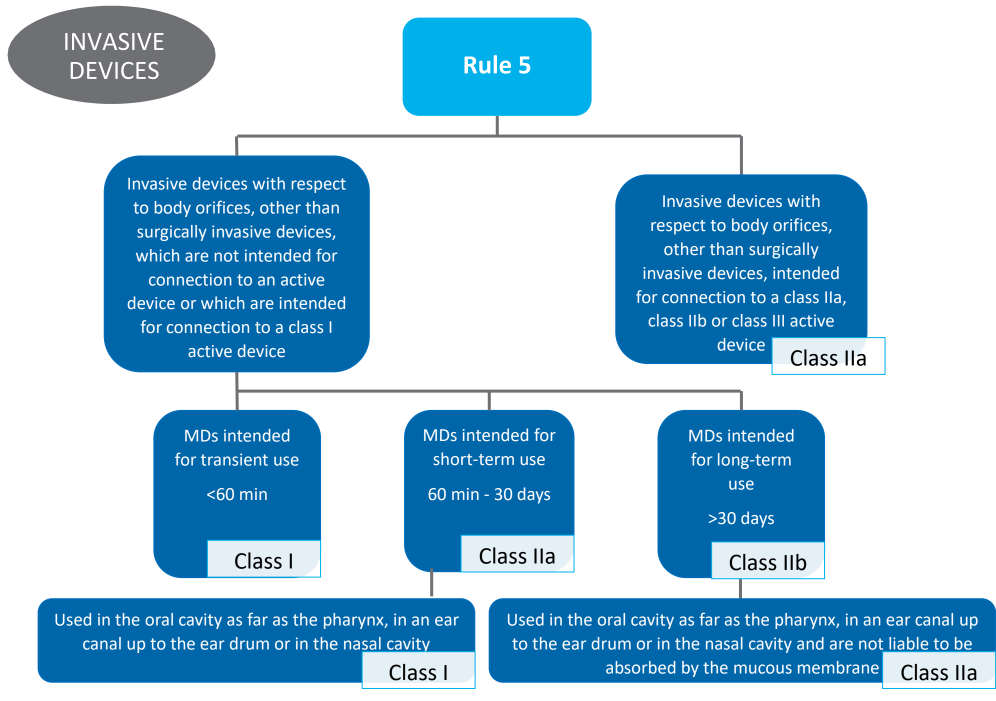


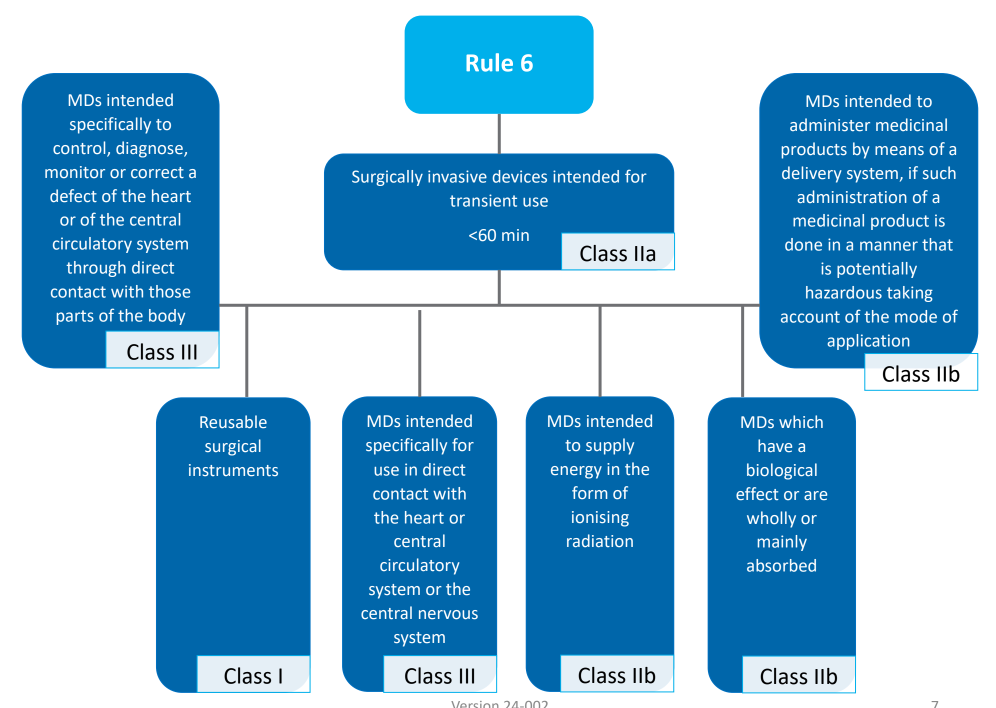


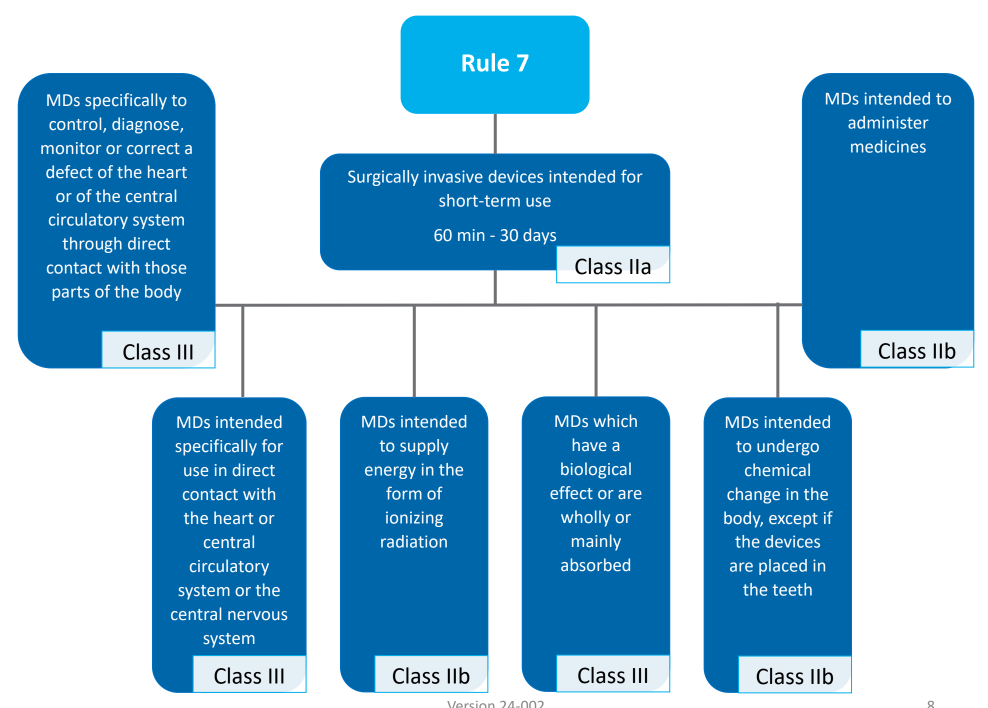


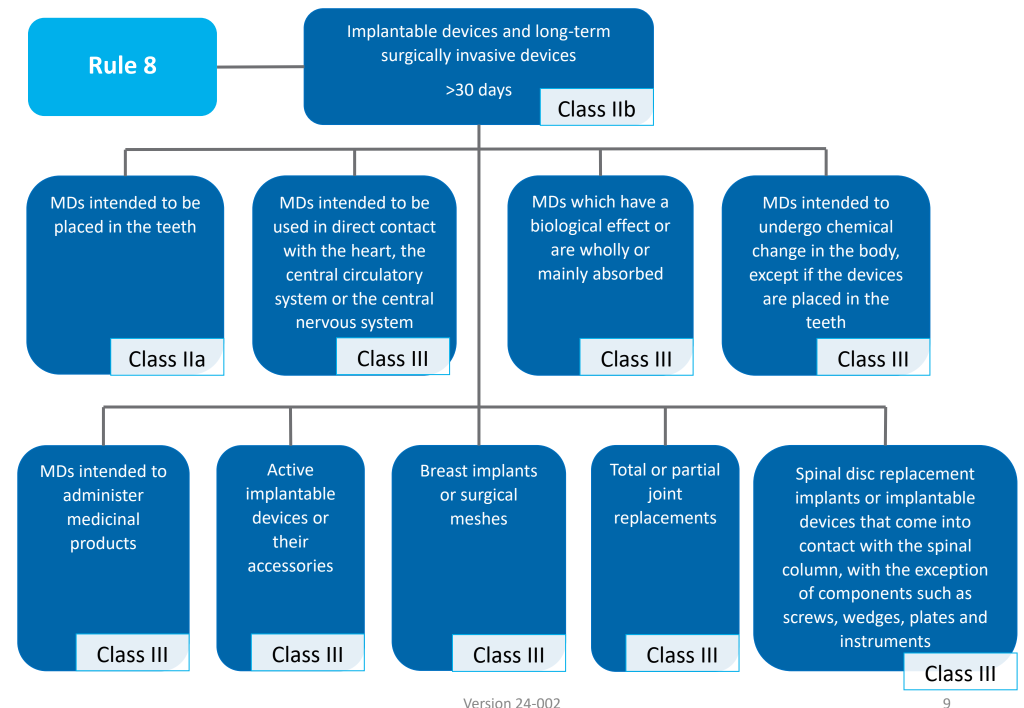












ACTIVE DEVICES

Rule 9

Active therapeutic devices intended to administer or exchange energy

Class IIa

MD's characteristics are such that they may administer energy to or exchange energy with the human body in a potentially hazardous way, taking account of the nature, the density and site of application of the energy

Active devices
intended to control
or monitor the
performance of
active therapeutic
class IIb devices, or
intended directly to
influence the
performance of such
devices

Class IIb

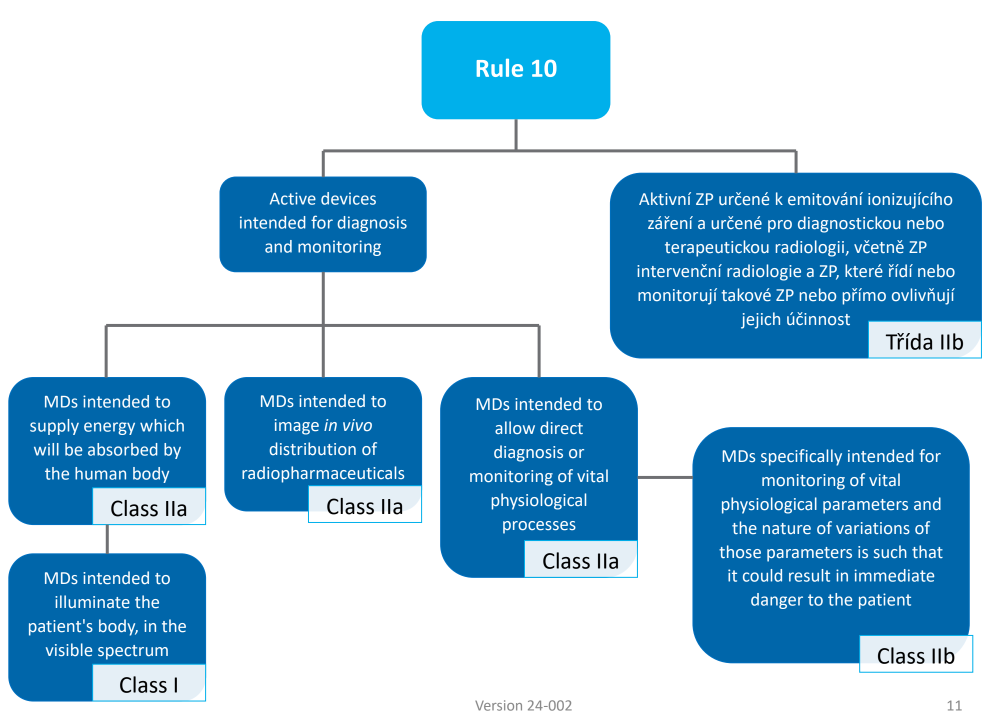
Active devices intended
to emit ionizing
radiation for
therapeutic purposes,
including devices which
control or monitor such
devices, or which
directly influence their
performance

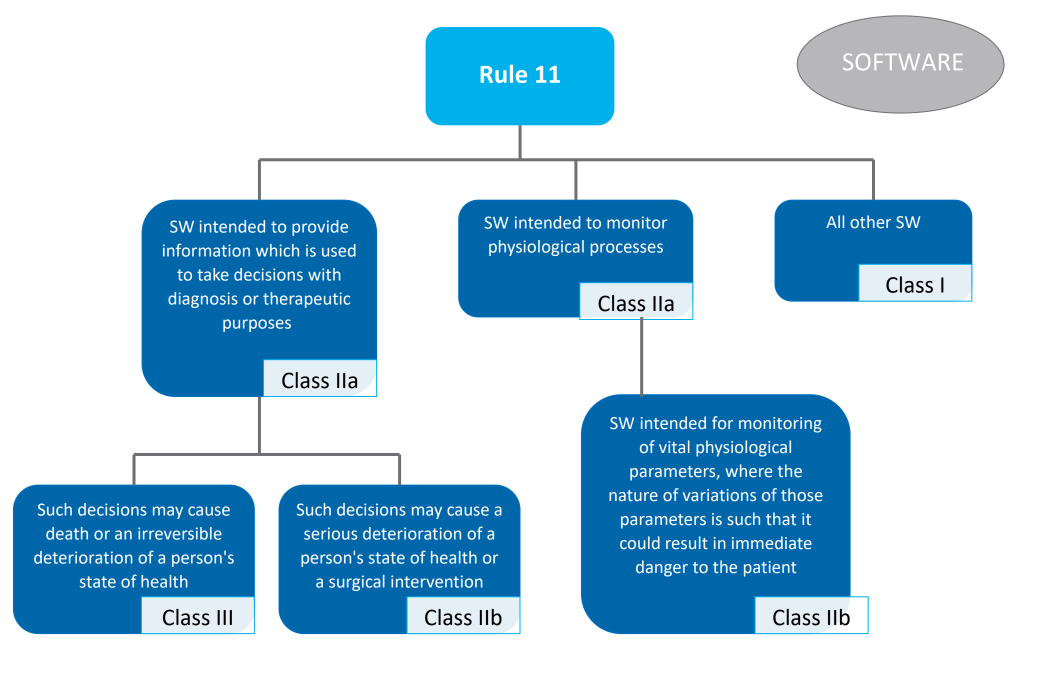
Class IIb

Active devices that are intended for controlling, monitoring or directly influencing the performance of active implantable devices

Class III

Class IIb







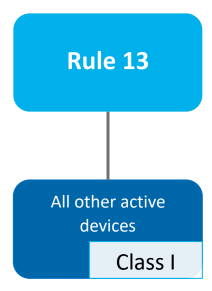
Active devices intended to administer and/or remove medicinal products, body liquids or other substances to or from the body

Class IIa

Active devices intended to administer and/or remove medicinal products, body liquids or other substances to or from the body, in a manner that is potentially hazardous, taking account of the nature of the substances involved, of the part of the body concerned and

of the mode of application

Třída IIb



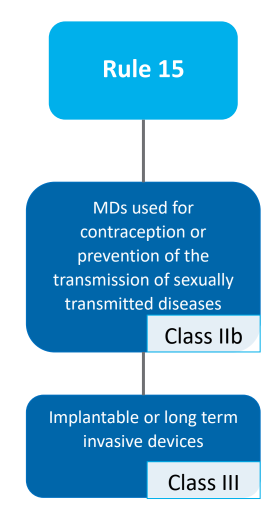
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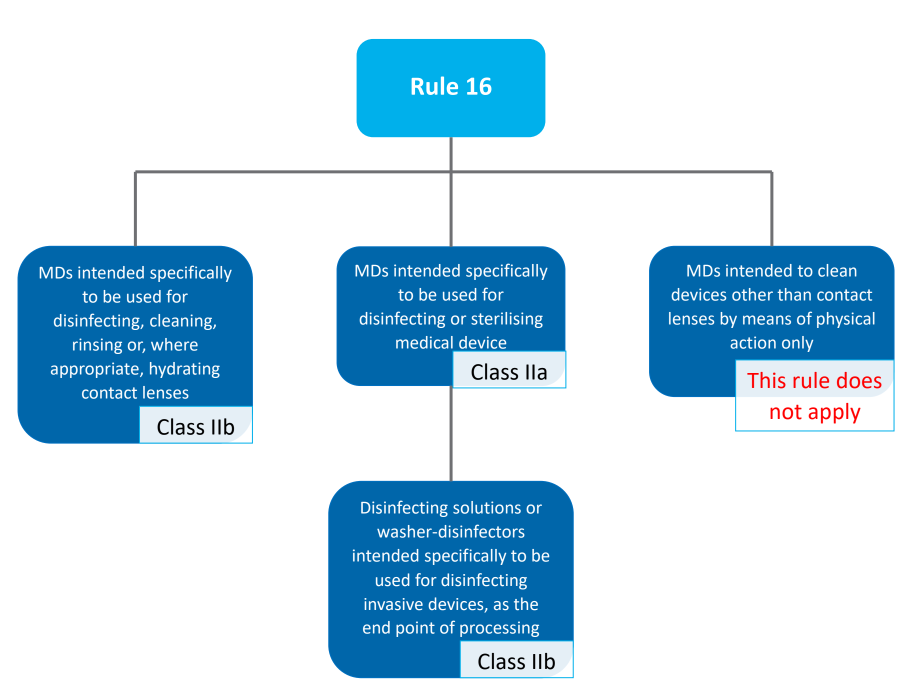
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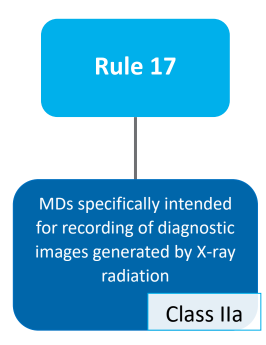


MDs incorporating, as an integral part, a substance which, if used separately, can be considered to be a medicinal product, as defined in point 2 of Article 1 of Directive 2001/83/EC, including a medicinal product derived from human blood or human plasma, as defined in point 10 of Article 1 of that Directive, and that has an action ancillary to that of the devices

Class III



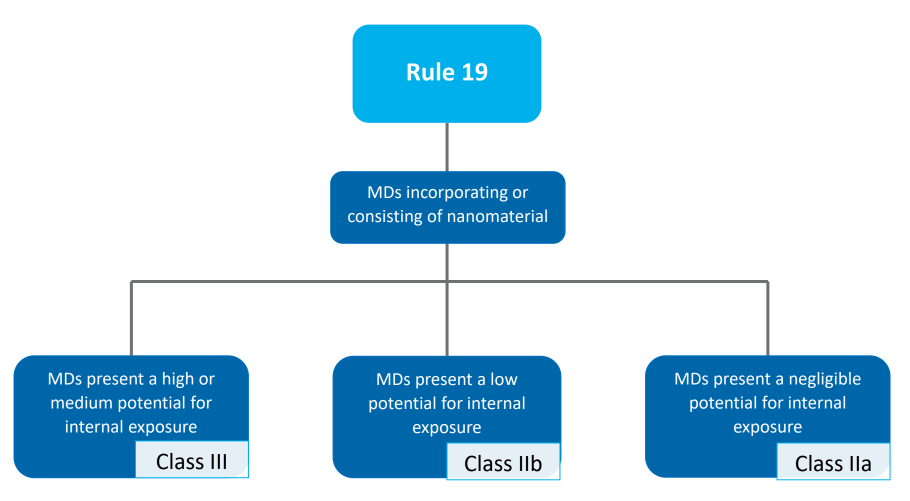




MDs manufactured utilising tissues or cells of animal origin, or their derivatives, which are non-viable or rendered non-viable and are not intended to come into contact with intact skin only, or manufactured utilising tissues or cells of human origin, or their derivatives, which are non-viable or rendered non-viable

Class III





Invasive devices with respect to body orifices, other than surgically invasive devices, which are intended to administer medicinal products by inhalation

Class IIa

Mode of action has an essential impact on the efficacy and safety of the administered medicinal product or they are intended to treat lifethreatening conditions

Class IIb

Devices that are composed of substances or of combinations of substances that are intended to be introduced into the human body via a body orifice or applied to the skin and that are absorbed by or locally dispersed in the human body

MDs or their products of metabolism, are systemically absorbed by the human body in order to achieve the intended purpose

Class III

MDs achieve their intended purpose in the stomach or lower gastrointestinal tract and they, or their products of metabolism, are systemically absorbed by the human body

Class III

MDs are applied to the skin or if they are applied in the nasal or oral cavity as far as the pharynx, and achieve their intended purpose on those cavities

Class IIa

All other cases

Class IIb

Active therapeutic devices with an integrated or incorporated diagnostic function which significantly determines the patient management by the device, such as closed loop systems or automated external defibrillators

Class III