










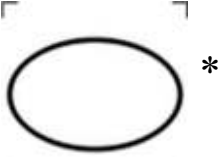
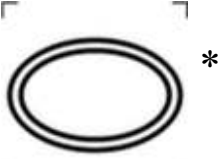
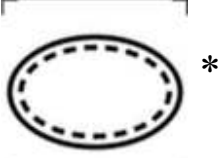
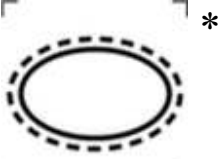




## List of Applied Symbols









As of: 07/2022

Symbol	Reg.-Nr ISO 7000	Title	Description
	3082	Manufacturer	Indicates the medical device manufacturer.
	From ISO 15223-1	Authorized representative in the European Community/ European Union	Indicates the authorized representative in the European Community/European Union.
	2497	Date of manufacture	Indicates the “date of manufacture”. The symbol shall be adjacent to the date that the product was manufactured, expressed as four digits for the year and two digits for the month and where appropriate, two digits for the day.
	2607	Use-by date	Indicates the date after which the medical device is not to be used.
	2492	Batch Code	Indicates the manufacturer’s batch code so that batch or lot can be identified.
	2493	Catalogue number	Indicates the manufacturer’s catalogue number so that the medical device can be identified.
	3725	Importer	Indicates the entity importing the medical device into the locale.







**DK 021 List of Applied Symbols**

Symbol	Reg.-Nr ISO 7000	Title	Description
	3724	Distributor	Indicates the entity distributing the medical device into the locale.
	-	Medical Device	Indicates the item is a Medical Device.
	2501	Sterilized using ethylene oxide	Indicates the method of sterilization using ethylene oxide.
	2503	Sterilized using steam or dry heat	Indicates the method of sterilization using steam or dry heat.
	3707	Single sterile barrier system	Indicates a single sterile barrier system.
	3704	Double sterile system	Indicates two sterile barrier systems.
	3708	Single sterile barrier system with protective packaging inside	Indicates a single sterile barrier system with protective packaging inside.
	3709	Single sterile barrier system with protective packaging outside	Indicates a single sterile barrier system with protective packaging outside.
	* A solid line identifies a sterile barrier system.		
	*The dashed line indicates a protective packaging that is designed to prevent damage to the contents or to help with aseptic		

## DK 021 List of Applied Symbols

Symbol	Reg.-Nr ISO 7000	Title	Description
			presentation. It does not provide a microbial barrier to maintain sterility.
			* This symbol shall be placed adjacent to or in combination with the symbol indicating the sterility.
	2608	Do not re-sterilize	Indicates a medical device that must not be sterilized again.
	2609	Non-sterile	Indicates that the device has not been sterilized.
	2606	Do not use if package is damaged and consult instructions for use	Indicates that a medical device that should not be used if the package has been damaged or opened and that the user should consult the instructions for use for additional information.
	0624	Store dry	Indicates that the medical device should be kept dry.
	0626	Store dry	Indicates that the medical device should be kept dry.
	0632	Temperature limit	Indicates the temperature limits to which the medical device can be safely exposed.
	1051	Do not re-use	Indicates a medical device that is intended for one use, or for use on a single patient during a single procedure.

## DK 021 List of Applied Symbols

Symbol	Reg.-Nr ISO 7000	Title	Description
	1641	Consult instructions for use or consult electronic instructions for use	Indicates the need for the user to consult the instructions for use.
	Derived from 2725	Contains or presence of natural rubber latex	Indicates the presence of natural rubber or dry natural rubber latex as a material of construction within the medical device or the packaging of a medical device.
	From prEN 15986	Contains the presence of phthalates	Symbol for use in the labelling of medical devices - Requirements for labelling of medical devices containing phthalates.
	From regulation (EU) 2017/745 or from directive 93/42/EWG	CE Marking of Conformity	Indicates manufacturer declaration that the product complies with the essential requirements of the relevant European health, safety and environmental protection legislation. XXXX indicates the identification number of the notified body.
	0434A	Caution	Indicates that caution is necessary when operating the device or control close to where the symbol is placed, or that the current situation needs operator awareness or operator action in order to avoid undesirable consequences.
	N/A	Unique device identifier	Indicates a carrier that contains unique device identifier information.

## DK 021 List of Applied Symbols

---

### Change history

Revision	Date	Created	Change content
1	01.07.2021	Anke Brenner	Addition of symbols to the sterile barrier systems and the importer.
2	04.05.2022	N. Froitzheim	Addition of symbols "Single sterile barrier system with protective packaging outside", UDI and Distributor. Adaption of description of some symbols in accordance to DIN EN ISO 15223-1: 2021 (E).