

Product Profile & Instructions for Use

Warwick House, Heathcote Way, Warwick CV34 6TE, United Kingdom T: 0044 (0)1926 422427 www.sasco.co.uk enquiries@sasco.co.uk

KD250 Kidney Dish

Class I Medical Device

Dimensions : 250 mm x 55 mm

Volume : 750 ml

Material : Blue Polypropylene

Case Size : 1 x 60



Intended Use

- Kidney dish, reusable.
- Kidney dish bowl to hold and transport medical/surgical instruments for a procedure. Also used for wound washing, and to hold vomit or oral secretions. Used in the treatment of disease.

Intended User

Product to be used by clinical, nursing, healthcare professionals, and decontamination technicians.

Product Features

- Autoclavable.
- Strong and durable.
- Guaranteed 1000 cycles.
- Light weight "about 50% lighter than stainless steel".
- Improved removal of residues.
- Smooth edges and surfaces eliminate paper rips during sterilization.
- Designed to maximise space for instruments and accessories.

Packaging Information

Catalogue	Case Bar Code	Primary	Secondary	Case Weight		
Number	(GTIN 14)	Packaging	Packaging			
KD250	15060098131118	Polythene bag 1 x 30	Cardboard case 2 x 30	6.6 Kg		

Product Symbols

Manufacturer	Catalogue number	EU Compliance	Medical Device	Authorized EC representative	Caution	Batch code	Use- by date	Non- sterile	Absence of natural rubber	Do not use if package is	Keep dry	Keep away from sunlight
***	REF	C€	MD	EC REP	\triangle	LOT	\square	NON	latex (LATEX)	damaged		*

Additional Information

- Harmonised System Product Tariff / Commodity Export Code 39.26.9097.
- **Country of Origin United Kingdom.**

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Manufacturer



- Warwick SASCo Ltd. Warwick House, Heathcote Way, CV34 6TE, United Kingdom.
- > Telephone: 0044 (0)1926 422427 E-mail: enquiries@sasco.co.uk

European Representative



- Medical Device Management Ltd. Block B, The Crescent Building, Santry, Dublin, D09 C6X8, Eire.
- Telephone: +353(0)18934143 E-mail: EU-REP@medicaldevicemanagement.com

Distributor/Importer

Standards

BS 5452:1977 Specification for hospital hollow-ware made of plastic Product complies with:

BS EN ISO 17664:2017 Processing of health care products – Information

to be provided by the medical device manufacturer.

BS EN ISO 20417:2021 Medical devices. Information to be supplied by

the manufacturer.

BS EN ISO 15223-1:2016 Medical Devices. Symbols to be used with medical device labels, labelling and information to be provided.

Company compliant to: BS EN ISO 13485:2016 Medical Devices. Quality management systems.

Requirements for regulatory purposes.

BS EN ISO 14971:2019 Medical Devices. Application of risk

management.

Referenced standards: BS EN ISO 15883 Washer disinfectors. Requirements.

BS EN ISO 17665 Sterilization of health care products. Moist heat.

BS EN ISO 25424 Sterilization of health care products. Low temperature

steam and formaldehyde.

BS EN ISO 11135 Sterilization of health care products. Ethylene oxide.

Instructions for Use

Storage and Handling

- Do not drop.
- Store in original packaging.
- Store in clean and dry conditions.
- Store off the floor on a shelf or pallet.
- Shelf life 36 months.

Preparation

- Product supplied non-sterile.
- Inspect products before processing.
- Withdraw damaged products and report to manager.
- Rinse with warm water to remove any packaging residues.

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Product not used in conjunction with any other device.

Cleaning and Disinfection

- Position products individually, avoiding contact with other items.
- Position products to allow water penetration and drainage.
- Machine clean in washer disinfectors compliant with ISO 15883.
- Use approved detergents, enzymatic or biocide cleaners, suitable for cleaning medical devices and surgical instruments which are compatible with polypropylene material (see detergent manufacturers' guide).
- Use detergents and disinfectants with a range between 2pH and 13pH.
- Ultrasonic cleaning is permissible.
- Manual cleaning is permissible separate product components and clean individually.
- Rinse with clean water.
- Dry with unwoven towels.
- Do not use abrasive materials.

Inspection Maintenance and Incident Reporting

- Visually inspect product to identify damage.
- Discard damaged product.
- Any serious incident that has occurred during the use of this product should be reported to the manufacturer, and the Competent Authority of the territory in which the user is established.

- Sterilization wrapping sheets and pouches must be compatible with polypropylene material and suitable for the chosen method of sterilization.
- Be aware that some wrapping sheets and pouches shrink during sterilization.
- Pouches must be large enough to contain the product without stressing the seals.
- We advise using a pouch or bag that is 25% larger than the product.

Sterilization

- Moist Heat sterilization (Autoclave) at 134.6°C (+2°C) for a minimum of 2 minutes.
- EO Ethylene Oxide "Gas" sterilization at maximum temperature of 65°C.
- H2O2 Hydrogen Peroxide "Plasma" sterilization at maximum temperature of 65°C.
- Sterilization procedures and machines to comply with ISO 17655, ISO 25424, and ISO 11135.

Disposal

- End of life is determined by wear and tear, visual inspection, or damage to product.
- If product is contaminated treat as clinical waste, and dispose of in accordance with clinical waste procedures.

Warnings 🔼

- Do not use abrasive powders or metal brushes as they may cause surface damage.
- Do not exceed 137°C.
- Do not exceed 8 minutes.

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Important Information for Processing

- The information provided by Warwick SASCo is the company's official instructions for this
- It is the responsibility of the processor to ensure the processing is performed in accordance with this information and that personnel, machines and equipment comply with relevant standards for
- Any deviation from this information requires evaluation for the effectiveness and potential adverse consequences.
- Additional information available on request contact: enquiries@sasco.co.uk

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