

Leg bag

Conveen®

Strap



Instructions for use 23323352 Version 1

Intended purpose

and/or retention of any aetiology.

Indications

Warnings

may occur.

The urine bag is intended to passively collect urine.

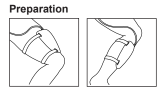
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Coloplast accepts no liability for any injury or loss that may arise if this product is used in a manner contrary to Coloplast's current recommendations.

Reusable products Straps should be replaced when the elasticity weakens.

How to use

Coloplast





The leg bag has an inlet tube which may be pre-attached to the bag or may be separate from the bag. The length of the tube determines where the bag can be placed on the leg.

If the tube does not have a pre-attached inlet connector or if the tube itself is not pre-attached to the bag you can shorten the tube to your preferred length. Simply cut the tube with scissors.

When cutting a corrugated tube, leave as much of the straight section as possible in place. This will ensure a secure connection between tube and connector when mounting the connector.

Attach the connector to the tube by pushing the thin end of the connector into the tube as far as possible.



TTTT-

Make sure the outlet tap is closed.

Attach the straps to the bag. Fit the bag

to your leg. Any surplus strap may be trimmed off.

Conveen siliconized strap

Use the longer strap at the top of the bag. If fitting the bag to the calf, wrap the top strap in a figure-of-eight behind and over the knee.

Reprocessing, washing, disinfection and/or (re)sterilisation may compromise product characteristics, causing additional risk of physical harm or infection to the user.

Urine bags are indicated for patients suffering from urinary incontinence

Re-use of the single use product is not recommended cross contamination

This leg bag is intended to be used while out of bed and moving about. Change to a bedside drainage bag with higher capacity while in bed and during sleep. Alternatively, leave your leg bag in place and attach a bedside drainage bag to expand the capacity of the overall urine collection system.

This product contains phthalates and should neither be used by children nor by pregnant or nursing women without consulting a health care professional.

Information

The bag is designed for single use.

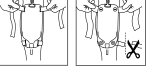
The product has a capacity of 600/800 mL.

The printed volume indicator is only an indicative volumetric measurement scale.

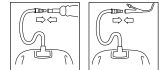
The straps can be washed at temperature of 40°C/104°F or lower.

The products are intended to be stored in the box until removed for use, as the box protects the products and the label on the box contains important information like batch code, use-by date and a unique barcode.

The device does not have to be removed before an MR scan as it is MR safe.



Application





If the connector has a protective cap, remove it.

Attach the connector to the urisheath or catheter.

The urinary collection system is now functional.



We recommend emptying the bag when it is a little over half full to ensure optimal drainage.

Removal

Disconnect the bag from the urisheath or catheter.

Unfasten the straps and remove the bag from your leg.





Change the bag according to local guidelines and recommendations.

The straps may be reused.

Empty the bag before discarding it.

The product is intended for single use only and should be disposed of in ac-cordance with local guidelines, e.g. with normal household waste.

Do not flush the product down the toilet.

Reporting of incidents If, during the use of this device or as a result of its use, a serious incident has occurred, please report it to the manufacturer and to your national authority.

Explanation of symbols

MD	Medical device
CE	Indicates that the product is in compliance with European legislation for medical devices
REF	Catalogue number
\leq	Use-by date (YYYY-MM-DD)
LOT	Batch code
$\sim \sim$	Date of manufacture (YYYY-MM-DD)
	Manufacturer
i	Consult instructions for use
(2)	Do not re-use
NON	Non-sterile
UDI	Indicates a barcode as containing Unique Device Identification
GTIN	Global Trade Item Number
×	Keep away from sunlight
	Recyclable packaging