Magnetic Drape
(PS4121 & PS4122)

EN

DIRECTIONS FOR USE

The following information should be read before using this device.

Indications:
The Purple Surgical Magnetic Drape minimises the risk of accidental injuries by providing an alternative to hand to hand transfer of surgical instruments. The Magnetic Drape is used to secure & exchange metal instruments on a sterile field.

Contraindications:
The Magnetic Drape should not be used on a patient with a pacemaker, or similar device, without first checking compatibility with the device manufacturer.

Device Description:
The Magnetic Drape is a single use product, and is supplied sterile
A lightweight, single-use magnetic drape that provides a secure 'hands free' transfer zone for metallic instruments, thus reducing the risk of third party inflicted sharps injuries.
Strong magnetic surface ensures secure retention of metallic instruments
Convenient single-use device
Lightweight construction makes the device particularly suitable for young or elderly patients
Waterproof construction minimises the risk of any pathogen cross infection
Available in two sizes

Colour: Purple
Dimensions: (Part Code: 4121) 25 x 40cm
(Part Code: 4122) 40 x 50cm

Directions For Use:

- The Magnetic Drape should be placed close to the surgical site.

- Instruments should be placed on, and retrieved from, the Magnetic Drape thus preventing the necessity of hand to hand transfer.

- After single patient use, the device is to be immediately disposed of as controlled medical waste.

Precautions/Warnings

⚠️ WARNING Hazard from a magnetic field

Use of the following substances may perforate the Magnetic Drape:

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- Oil, ointment and oil containing soaps (green soap)
- Hydrocarbon solvents: ether, acetone, benzene, freon, alcohol or oxidizing agents.
- Sulphur, acetic and hydrochloric acids, lye and sodium hypochlorite (bleach).
- Phenol and hexachlorophene products.
- Quaternary compounds and benzalkonium chloride.

How Supplied

These device(s) are designed and sold for single use only as defined in Article 1 (n) of directive 2007/47/EC. As such re-processing and, or, re-sterilisation after initial use is not permitted.