



The following should be read thoroughly before use:

Important Notes:

This document is designed to provide instructions on how this product/s is used. It is not a reference to clinical techniques, and does not replace local or national guidance or protocols.

Intended User:

For use by both suitably trained healthcare professionals in a healthcare establishment and 'Lay Users' in a home use environment.

Intended Purpose:

Provides support and stabilization for the leg and knee.

Indications For Use:

- Post trauma
- Post surgery
- Immobilisation for pain relief
- Cast removal

Product Characteristics / Features:

Knee Immobiliser - available as either:

- Universal - Knee circ 36-41cm - Varying lengths
- 20" Length - Varying knee circs

Key functional elements include:

- 3 panels of soft foam laminated material for comfort and exact fit
- Straight lateral and medial aluminium stays provide increased stability
- Velcro closure for secure fit
- Detachable side panels can be angled for a customised fit

How it's supplied:

It is supplied non-sterile and ready for single patient use.

Devices Required For Use, And Interoperability

Requirements:

None

Storage Conditions:

Store in a cool and dry environment.

Care and Maintenance:

- Wash periodically by hand with a damp cloth and mild soap
- Use a towel to absorb most of the dampness and then allow to dry at room temperature.
- Do not expose to alcohol, ointments or solvents.
- The device should be safety checked by the providing service and/or user every 6 months.

Contraindications:

None

Precautions / Warnings / Residual Risks:

If allergic skin rashes occur, please consult your healthcare professional

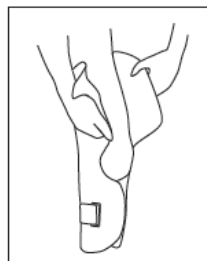
Possible Adverse Reactions:

None

Directions For Use / Fitting Instructions:

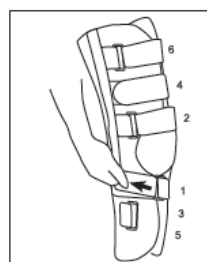


1. Refer to the sizing charts to make sure that you have the correct size.



2. Undo all the straps and remove the side panels.

3. Wrap the sleeve around the leg, making sure that the knee cap is visible through the opening.



4. Fasten the sleeve using the Velcro straps to ensure a snug fit around the leg.

5. Contour the side panels to shape of the leg by bending the aluminium stays and then reattach to the sleeve.

6. Fasten the straps through the D-rings in the order indicated.

Single Patient Use Precaution:

This device is designed and sold for single patient use only. The effects of any unauthorised reuse can result in the following complications:
 Risk of Cross contamination;
 Risk of mechanical fatigue, and/or associated failure.

Disposal

Discard after single patient use and dispose of as common waste according to national guidelines.

Serious Incident Reporting

Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

Sizes & REF / Codes

20" Tri Panel Knee Immobiliser

Size	Knee Joint Circ		Code
	Inches	CM	
Small	13 - 14	33 - 36	KI-20-S
Medium	14 - 15	36 - 38	KI-20-M
Large	15 - 16	38 - 41	KI-20-L
X Large	16 - 18	41 - 46	KI-20-XL

Universal Tri Panel Knee Immobiliser

Length	Knee Joint Circ		Code
	Inches	CM	
12"	14 - 16	36 - 41	KI-UNI-12
14"	14 - 16	36 - 41	KI-UNI-14
16"	14 - 16	36 - 41	KI-UNI-16
18"	14 - 16	36 - 41	KI-UNI-18
20"	14 - 16	36 - 41	KI-UNI-20
22"	14 - 16	36 - 41	KI-UNI-22
24"	14 - 16	36 - 41	KI-UNI-24

LABELLING SYMBOLOGY
 EN ISO 15223-1:2021, unless otherwise stated

	Manufacturer		Consult Instructions for Use		Unique Device Identifier
	Authorised representative in the European Community		Caution		Single Patient Use
	Medical Device		Non-Sterile		Quantity - Abbreviation
	Catalogue Number		CE Mark - EU Regulation 2017/745		Contains Latex
	Batch Code		UKCA Mark - UK Regulation SI 2002 No 618		Date of Manufacture

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