

Mesh Sling

Nursing Care

INTENDED USE

The **Mesh Sling** is a medical device which is intended to be used for the immobilization of the arm/shoulder by injuries or fractures.

COMPOSITION

Product composed of mesh and polyester fastening strips, with polypropylene rings.

INSTRUCTIONS FOR USE



- Deploy the immobilizer deploying the strips.
- Put the arm inside. Support the thumb and adjust the tape B.
- The patient's arm must rest on the same square.
- Pass the tape fastening A above the arm and adjust the measure through the ring.
- Pass the subject strap D around the chest, immobilizing the arm.

PRODUCT CHARACTERISTICS

Designed for easy application and adjustment. It has velcro system for closing and adjusting.

It supports the forearm and wrist. Incorporates support strip for the thumb.

Presents a vertical adjustable strap with hooks, its function is to distribute the weight of the arm across his back and shoulder for great comfort. The strap C facilities the adjustment of the arm sling.

This material gives to the product: lightness, breathability and comfort, without losing strength or adaptability to the body.

PRECAUTIONS AND RECOMMENDATIONS

The use by more than a patient may provoke a cross-infection.

Do not use the product on injured skin.

Do not use the product if it is polluted, dirty or evidently deteriorated.

STORAGE AND SHELF-LIFE

Keep away from sunlight. Keep dry. Shelf-life: 5 years.

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PRODUCT DIMMENSIONS

In order to choose the correct size, measure the length of the forearm from the elbow to the knuckles.



REFERENCE	MODEL	L (cm)	W (cm)	L. Strap (cm)	W. Strap (cm)
1012825	L	46	21	Chest: 111 Neck: 97	4 4
1012835	M	40	19	Chest: 108 Neck: 97	4 4
1012845	S	32	16	Chest: 74 Neck: 74	4 4

CONFIGURATION							
REFERENCIA	UNITIES BOX	MEASUREMENTS BOX (cm)	WEIGHT BOX* (kg)				
1012825	10 un/box	21x15x15	0.59				
1012835	10 un/box	21x15x15	0.57				
1012845	10 un/box	21x15x15	0.45				

* TOLERANCE: Weight ± 0.2 Kg

IN COMPLIANCE WITH:

- Directive 93/42/EEC of the Council, Directive 2007/47/EC of the European Parliament and of the Council and Regulation (EU) 2017/745 of the European Parliament and of the Council.
- Class I, non-sterile (according to Annex IX of Directive 93/42 and Annex VIII of Regulation 745/2017).
- Enterprise certified by ISO 13485 and ISO 9001.
- Medical Devices Manufacturer Operating License No.: 3689-PS.







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