







MORTUARY BODY SLING

This guide covers all products with the prefix MSS in the product code.

The **Mortuary Body Sling** is for lifting and lowering bariatric deceased safely and easily.

USER INSTRUCTIONS



	<h1>1</h1> <p>Before each use, check each strap for signs of wear or damage.</p>
<p>Slide the straps under the deceased in line with the hooks on your overhead lifting system; this is generally under the deceased person's head, shoulders, waist, legs and feet area.</p>	 <h1>2</h1>
	<h1>3</h1> <p>Connect the straps to the hooks on the overhead system. Check they are all secure.</p>
<p>Lift the deceased and then lower gently into the coffin, or transfer from one surface to another. Remove the straps by pulling each out from one side.</p>	 <h1>4</h1>
	<h1>5</h1> <p>The slings can then be wiped clean or laundered ready to be used again.</p>

MATERIAL CONTENT

Fabric: 100% Polyester Webbing

STORAGE AND HANDLING

- Store in a clean dry environment that is not subject to excessive heat.
- Keep away from sunlight.
- Store flat and avoid placing heavy objects on top of the product.

WARNINGS AND PRECAUTIONS

- Do not leave the on the floor when not being used.
- Inspect the product before use for signs of damage or tears.

CLEANING INSTRUCTIONS

- Wash as per laundry instructions.
- Can be wiped clean with natural water and detergent or wipes that don't contain alcohol, solvents, bleaching or abrasive agents.

DISPOSAL INSTRUCTIONS

- Non-recyclable.
- Dispose of as clinical waste if it has been used by an infectious person.
- Otherwise dispose of through normal waste management.



Keep dry



Keep away from sunlight



Caution



Do not use if package is damaged



Storage Temperature Limit



Wash at 30-74°C. Tumble dry on low heat. Do not iron. Do not use bleach. Do not use fabric conditioner



Patient Information website



Medical Device



MANUFACTURER

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EU REPRESENTATIVE: Obelis S.A., Bd Général Wahis 53, 1030 Brussels, Belgium



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HOSPITAL DIRECT (HD) ADDITIONAL SYMBOLS

At HD we understand that not everyone is familiar with all equipment, so to help therapists and healthcare professionals in assessing a product's suitability for a client, we have included two assessment tools, Functional Independence Measure (FIM) and Easy Guide Symbols. These will enable the healthcare professional to assess their client and decide whether or not a product will help the person based on their level of dependence and functional ability.

FUNCTIONAL INDEPENDENCE MEASURE (FIM) ASSESSMENT

Our FIM assessment guide for each product will help you decide the suitability of this product for the person's ability and need. Based on the standard criteria from Level 6 where the person can use the product unaided and unsupervised to Level 1 where all the assistance is provided by the carer and the client can do nothing, this guide is easy, quick and simple to use to check suitability against ability and circumstance.

FIM LEVELS QUICK GUIDE. SUBJECT VS CARER HELP



EASY GUIDE SYMBOLS

We use four symbols to indicate where a product is suitable and safe for the person to use unassisted, with a single carer or with multiple carers. These symbols indicate the minimum recommendation.



HD has a policy of continuous development and, as such, reserves the right to alter specifications (including measurements, materials and colours) without prior notice. If any serious incident has occurred in relation to the device it should be reported to the manufacturer and the competent authority in which the user and/or patient is established.