

BONEFOAM™

RESTORE

DUAL LEG SUPPORT



Restore | Dual Leg Support is a nonsterile, single-patient use, therapeutic positioner intended to align the legs with the feet, ankles and knees supported in a neutral position to prevent external hip rotation in the supine position.

Healthcare Facilities:
To reorder the following product in bulk,
please contact sales@bonefoam.com

• 737.RDLS - Restore | Dual Leg Support



BoneFoam, Inc.
20175 County Road 50
Corcoran, MN 55340
USA

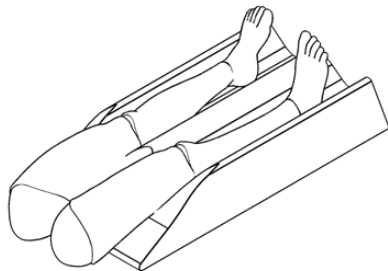
Email: customerservice@bonefoam.com
Phone: (763) 559-1830
Toll-free: 1 (877) 861-2663
Fax: (763) 559-1822

bonefoam.com

RDLS-eIFU Rev.0 2025-11-04

INSTRUCTIONS FOR USE

Talk with your physician or therapist for recommendations on how Restore | Dual Leg Support can be incorporated into your recovery or sleep routine.



WARNINGS/PRECAUTIONS/CONTRAINDICATIONS

- Intended to contact intact skin only.
- Risk of cross contamination if used by multiple patients.
- Incorrect positioning of the patient with this product can increase the risk of:
 - Pressure injury resulting in markings or ulcers
 - Peripheral nerve injury, which can lead to temporary or permanent paralysis
 - Venous pressure, which can lead to ischemia
 - Instability, which can lead to patient falls
- Not intended for patient transport.
- Ensure device is compatible with work surface, procedural instruments, and equipment.
- Risk of fire. Below are flammability ratings of the materials:
 - Foam: Rated either UL 94 HBF or CAL 117

CARE, HANDLING AND USEFUL LIFE

- Avoid using the product if package is damaged during transportation.
- Transport and store in dry conditions and out of direct light where possible.
- Product can be cleaned with a dry cloth if needed.
- To prevent patient and/or user injury and/or equipment damage, examine the product for potential damage or wear prior to use. Do not use the product if damage is visible, if parts are missing, or if it does not function as expected.
- Avoid stacking items on top of product when storing.
- Do not submerge the product in water. Equipment damage can occur.
- Prior to using this product, read the Instructions for Use.
- Discontinue use if foam has degraded or has lost resilience.
- Compression packaged foam may require up to 10 minutes before functionally usable and up to 24 hours before reaching final dimensions.

SAFE DISPOSAL INSTRUCTIONS

Customers should adhere to all federal, state, regional, and/or local laws and regulations as it pertains to the safe disposal of medical devices and accessories.








PACKAGING






The product is packaged in a clear polythene bag inside a corrugated cardboard box. Components may be compressed to minimize packaging and storage footprint.

SAFETY INFORMATION

The techniques detailed in this manual are only manufacturer's suggestions. The final responsibility for patient care with respect to this product remains with the attending physician. Any serious incident that has occurred in relation to this product should be reported to the manufacturer and regulatory authority, where required. Keep this manual available for future reference.

SKU	DESCRIPTION	INTENDED USE	INDICATIONS
INDIVIDUAL			
737.RDLS	Restore Dual Leg Support	Nonsterile, single-patient use, therapeutic positioner intended to align the legs with the feet, ankles and knees supported in a neutral position to prevent external hip rotation in the supine position.	Indicated for use by healthcare professionals and patients to align and support the patient's legs during sleep and recovery.

SYMBOL	DESCRIPTION
	Indicates the device is a medical device
	Indicates the medical device manufacturer
	Indicates the country where the medical device was manufactured and date of manufacture
	Indicates the manufacturer's lot number
	Indicates the medical device Global Trade Item Number
	Indicates the manufacturer's catalogue number
	Indicates the unique product identifier information for the device

SYMBOL	DESCRIPTION
	Indicates the need for the user to consult the instruction for use
	Indicates the device does not contain natural rubber or dry natural rubber latex
	Indicates the device is single-patient use
	Indicates a medical device should not be used if the package has been damaged or opened
	Indicates quantity in package