

Foot Garment RFG210

Universal size

Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.

DVT System

Instructions for use

Secure the pump at the foot of the bed or in place of use. Attach the tube set to the pump by pushing in the silver latch on the female pump connector and then inserting the male tube connector into the pump connector until a click is heard.

Garments may be used on either foot. Unfold the garment(s) and position the inflatable center bladder directly below the arch of the patient's foot.

SNUGLY wrap the garment(s) around the patient's foot and secure the hook tabs. Wrap the ankle straps around the back of the ankle and on top the foot.

25°

Attach the garment(s) to the tube set by pushing in the silver latch on the female tube connector and inserting the male garment connector firmly into the tube connector until a click is heard.

Ensure the pump is held firmly onto the bed frame. Foot garments should be removed prior to patient ambulation or transportation.

Turn the pump on by pressing the POWER button. If the LOW BATTERY light is solid red and producing a constant audible alarm, the battery will need to be charged. To charge the unit, plug the supplied power adaptor into a power outlet and then plug the round connector at the end of the cord into the top of the pump. The BATTERY CHARGE light will flash green

while the unit is charging. The unit may be used while it is being charged. When fully charged, after 16-48 hours, an audible double-beep is presented once and the BATTERY CHARGE light (green) will be lit constantly.

Ensure the TYPE light indicates FOOT GARMENT. If not, press the TYPE button until the light is illuminated.

Ensure the MODE light is not lit for SINGLE GARMENT. If only one garment is needed, press the MODE button to select SINGLE GARMENT.

The pump is now ready for use. Refer to the user operating manual for complete information on the use of the system.

Indications

Intermittent Pneumatic Compression (IPC) is indicated for use for the prevention of deep vein thrombosis due to the presence of risk factors for thrombus formation during orthopedic, trauma, urologic, neurology, critical care, general medicine, obstetrics, and general surgery.

Recommendations

Garments should be removed regularly to inspect the skin.

Patients should be instructed in the proper use of the system and report any problems to the nursing staff.

The system should be used continuously until the person is fully ambulatory. Foot garments should be removed before ambulating.

If the garment cannot be applied to a limb during surgery, it may be applied to the limb once the patient reaches the recovery room. For a non-surgical patient, the system should be applied as soon as the risk of DVT is identified.

Contraindications

The Restep® system should not be used in the following conditions:

- Severe arteriosclerosis or active infection.
- Suspected or known acute DVT
- Severe congestive heart failure or any condition where increased fluid to the heart may be detrimental.
- · Existing pulmonary edema.
- Local skin or tissue conditions in which the garments would interfere.

If in doubt, refer to the patient's physician before using device.

Cautions

Garments should be removed immediately if the patient experiences any unexplained sensations, numbness, or pain.

When used for DVT prophylaxis, continuous use is recommended and any prolonged interruption of therapy should be done in consultation with the patient's physician.

Garments are single-patient use, non-sterile, and latex-free. Garment bladders are PVC-free.



Pack contains one Single Compression Sleeve Non-Sterile (Devices HLD Processed)

Sustainability Solutions

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NOT Reuse





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Pump RSP101 only.

For use with Restep

Made in the Dominican Republic

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Restep DVT System

Warranty

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STRYKER SHALL NOT BE LIABLE FOR ANY DAMAGES TO THE EXTENT CAUSED BY ANY DEFECT IN MATERIAL, WORKMANSHIP OR DESIGN BY THE ORIGINAL MANUFACTURER OF THE PRODUCT OR ANY ACT OR OMISSION OF THE ORIGINAL MANUFACTURER OF THE PRODUCT.

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If a valid warranty claim is received within thirty (30) days of the expiration of the applicable warranty period, Stryker will, in its sole discretion: (1) replace the product at no charge with a product that is at least functionally equivalent to the original product or (2) refund the purchase price of the product. If a refund is provided by Stryker, the product for which the refund is provided must be returned to Stryker and will become Stryker's property. In any event, Stryker's liability for breach of warranty shall be limited to the replacement value of the defective or non-conforming part or component.

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