

Reprocessed by



HEALTH SERVICES

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Instructions for Use
Hygia Reprocessed Aircast Venaflo Elite Foot Cuff IFU
Reprocessed Device for Single Use.

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

NON-STERILE NOT MADE WITH NATURAL RUBBER LATEX

Sleeves Description

Hygia Reprocessed Aircast Venaflo Elite Foot Cuff

Indications for Use

The Hygia Health Services Reprocessed Aircast foot cuffs are designed to increase venous blood flow in at-risk patients in order to prevent deep vein thrombosis. The cuffs are a non-invasive therapeutic method for prevention of deep vein thrombosis. The cuffs are also used in the treatment of venous leg/foot ulcers and edema which are disorders associated with venous insufficiency.

Warnings/Precautions

- Proper foot cuff positioning must be assured to avoid the potential for pressure points on the foot.
- Ensure proper connections to the Venaflo pump and cuffs.
- Kinked or twisted tubing may restrict air flow.
- Foot cuffs should be removed if patient experiences numbness, tingling, or foot/leg pain.
- To minimize local air movement when using foot cuffs in operating room, cuff cooling should be turned off, if available.

Directions for Use

1. Remove Aircast foot cuffs from plastic bag.
2. Apply each cuff with the aircell centered on bottom of foot using diagram on material of foot cuff.
3. Secure foot cuff straps snug, but not tight.
4. Be sure to check that all tube assembly connections are tight and that cuffs are snug to achieve full pressure. Always turn off pump before readjusting or removing device.
5. Refer to original equipment manufacturer manual for further instructions.

Contraindications

1. Any local leg condition in which sleeves/cuffs would interfere such as dermatitis, vein ligation (immediate postoperative), gangrene, or recent skin graft
2. Severe arteriosclerosis or other ischemic vascular disease
3. Massive edema of legs or pulmonary edema from congestive heart failure
4. Extreme deformity of leg
5. Suspected existing deep venous thrombosis
6. Congestive heart failure

The OEM information listed on the label is provided as device ID prior to reprocessing and may contain the trademarks of unrelated third parties that do not sponsor this device.

This product and its packaging have been exposed to ethylene oxide gas (EtO). Even though the product is processed in compliance with all applicable laws and regulations relating to EtO exposure, proposition 65, a State of California voter initiative, requires the following notice:

Warning: this product and its packaging have been exposed to ethylene oxide. The packaging may expose you to ethylene oxide, a chemical known to the State of California to cause cancer or birth defects or other reproductive harm.