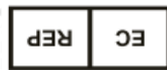




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**NORTH AMERICAN RESCUE®**



REF 30-0071 BOA XL

REF 30-0009 NSN: 6515-01-537-2611

# BOA®-Constricting IV Band

**WARRANTY:** The product or components contained in this package may constitute a medical device for which specific training is required for proper use. North American Rescue, LLC. (NAR), warrants that the product is merchantable and fit for its specified purpose. NAR expressly disclaims all other express or implied warranties relating to the product; any use beyond the product's specified purpose; or use by any party who is not trained or legally authorized to use such product. Use only as directed by your EMS authority or under the supervision of a physician.

**DISPOSAL:** The BOA® is a single use device and designed for disposal after use. Do not attempt to clean or reuse the device, as it may increase the possibility of cross contamination. Dispose of the device in a manner ensuring the isolation of potential substances in accordance with universal protocols. Contact NAR for more information as needed.



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REV02132020

PN: ZZ-0087

requirements.

Within the European Union (EU), the BOA® Constricting IV Band is not indicated for pediatrics and pregnant women due to EU regulatory

## Contraindications

The BOA® should be used only by trained medical professionals. Improper use could result in injury to casualty. Reuse of this device will degrade efficacy, resulting in adverse casualty reaction, including potential death. Continually monitor casualty to ensure BOA® is functioning per medical protocols. In the event of a BOA® malfunction, follow local protocols and report any serious incident to North American Rescue or authorized representative and the competent authority of the Member State.

## Warning

The BOA® Constricting IV Band provides circumferential pressure on either of the upper extremities for the purpose of obtaining peripheral venous access.

## Intended Use



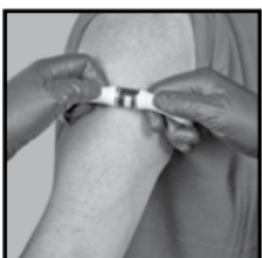
# Instructions for Use



1. To find the correct size, place the BOA<sup>®</sup> (in its relaxed state) gently around the arm. It should measure 1/2 to 2/3 around the extremity.



2. Stretch BOA<sup>®</sup> straight out and place as high as possible around upper arm.



3. Keep fingers underneath the connectors as you secure in place to prevent pinching of the skin.



4. Roll BOA<sup>®</sup> gradually down the arm. Insert IV/Saline Lock in accordance with your protocol.



5. Hold inserted IV/Saline Lock with one hand, press the "Quick Release" button and pull to remove the BOA<sup>®</sup>.
6. Monitor casualties with compromising clinical conditions. Evacuate casualty

to secondary treatment facility, advising follow on care to monitor casualty for allergic reactions or infections.

7. In the event of device issues refer to medical protocols.

## Helpful Hint:

*Hold the female end containing the "Quick Release" in the non-dominant hand when applying the BOA<sup>®</sup>. This is the hand you are going to use when releasing the BOA<sup>®</sup> while the dominant hand holds the IV/Saline Lock in place*

## Harmonized Standard Symbols:

- LOT Lot number
- Single use
- Do not resterilize
- Consult instructions for use
- Manufacturer
- Authorized Representative
- Device part number
- Not for use in MRI
- Date of manufacture