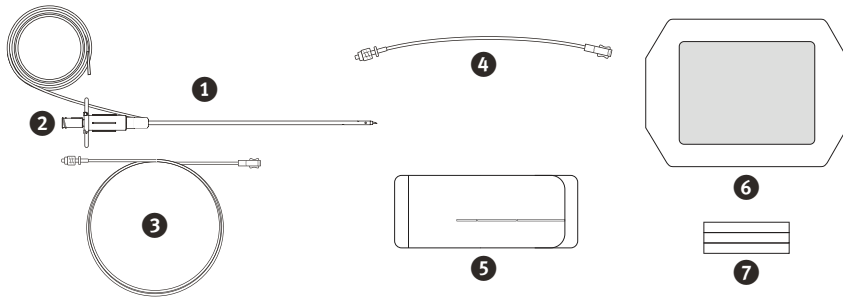


ON-Q[®] PAIN RELIEF SYSTEM QUIKBLOC[®] OVER-THE-NEEDLE CATHETER SET



Contents

- 1 QUIKBLOC[®] Over-the-Needle Catheter Set (some models include an integrated stimulating cable)
- 2 Removable Needle Wing
- 3 Non-DEHP 24 in. Needle Extension Set
- 4 Non-DEHP 6 in. Catheter Extension Set
- 5 Connector Securement Device
- 6 Occlusive Dressing
- 7 Adhesive Strips

DESCRIPTION

The ON-Q[®] QUIKBLOC[®] Over-the-Needle Catheter Set consists of a catheter assembled over an echogenic needle.

- Needle: 20 GA ECHOBRIGHT[®] Echogenic
- (Stimulating and non-Stimulating)
- Catheter: 16 GA open tip with lateral ports and 10 mm depth marks

⚠️ WARNINGS

- **Avoid placing the catheter in joint spaces. Although there is no definitive established causal relationship, some literature has shown a possible association between continuous intra-articular infusions (particularly with bupivacaine) and the subsequent development of chondrolysis.**
- **Do not reinsert the needle through the catheter, after the catheter is placed. This may damage the catheter and result in the catheter breaking off inside the patient.**
- **Ensure that needle or catheter is not placed in a vein or artery. Inadvertent intravascular delivery may result in systemic toxic effects. Refer to the drug manufacturer's package insert.**
 - **If blood is observed during placement of the needle or catheter, it may have been placed incorrectly. Stop immediately and follow currently accepted medical practices.**
- **Do not bend the QUIKBLOC[®] Set to excess as this may cause product damage.**
- **Consider location of needle and catheter insertion to avoid injury to nerves, blood vessels, organs and other anatomical structures.**
- **Catheter should not be cut as this may lead to failure of catheter function, and/or retention of catheter components on removal.**

⚠️ CAUTIONS

- **STERILE** Product is ethylene oxide sterilized.
- Do not use if package has been opened or is damaged.
- **MR** Catheter is MR safe.
- Single use only.
- Do not resterilize or reuse. Reuse of the device could result in the following risks:
 - Damage to the catheter and needle.
 - Increased risk of infection.
- Consult standard textbooks and practice guidelines for specific techniques and cautions for continuous peripheral nerve block procedures.
- Maintain catheter per standard hospital protocol.
- To help prevent tubing misconnections, a Catheter Site Identification Label is provided. Information such as the site and route of administration may be written on the label and then affixed to the catheter.
- If the catheter disconnects or a leak occurs at the catheter connection, close the clamp on the infusion system and discontinue use.

INDICATIONS FOR USE

The ON-Q[®] QUIKBLOC[®] Over-the-Needle Catheter Set is indicated for delivery of medication for regional anesthesia and pain management. Route of administration may be intraoperative, percutaneous, or perineural.

CONTRAINDICATIONS

The ON-Q[®] QUIKBLOC[®] Over-the-Needle Catheter Set is not indicated for epidural and intravascular delivery.

For Customer Service, please call +800.448.3569 (USA only) and Outside the USA, 1-949-923-2400 or visit www.halyardhealth.com for the latest product information and Technical Bulletins.

Instructions For Use

USE ASEPTIC TECHNIQUE PLACING THE CATHETER

1. Prime and connect the 24 inch injection set to the QUIKBLOC[®] Set to facilitate medication administration (optional).
2. Prepare the procedure site according to your facility's protocol.
3. Introduce the QUIKBLOC[®] Set through the skin.
 - Note:** Using an 18 GA needle to create an entry point can help ease the insertion process.
 - Note:** The ↑ on the hub of the needle aligns with the needle bevel facing upward. The arrow may be used as a reference point on QUIKBLOC[®] catheter position during placement.
4. Gradually advance the QUIKBLOC[®] Set to the desired location using ultrasound and/or nerve stimulation techniques.
 - ⚠️ **Caution:** If excessive resistance is encountered, stop, remove the QUIKBLOC[®] Set and restart the procedure.
5. Aspirate and inject medication as needed.
6. Once the desired location is confirmed, stabilize the catheter at the hub and remove the needle, leaving the catheter in place.
 - ⚠️ **WARNING: To avoid damaging the catheter, never reinsert the needle through the catheter after the catheter is placed.**
7. Attach the 6 inch extension set to the catheter hub.
8. Secure the catheter hub in place with the Connector Securement Device.
 - Note:** For best adhesion, prepare and dry skin. See reverse side for illustrated instructions.
 - a. Lift the top flaps and remove the paper backing to expose the adhesive strip.
 - b. Place catheter hub on bottom adhesive strip and secure with the top flaps over the catheter hub.
 - c. Peel the paper backing from the Connector Securement Device and secure in desired position on the skin.
9. Affix the catheter identification label to the extension set.
10. Apply adhesive strips, occlusive dressing or other securement devices.
11. Connect the extension set of the catheter to an infusion device. Ensure the connection is secure.

CATHETER REMOVAL

1. Remove dressing and any securement devices from the catheter site.
2. Grasp the catheter close to the skin and gently pull to remove. The catheter should be easy to remove. Do not tug or quickly pull on catheter during removal.
 - ⚠️ **Cautions:**
 - If resistance is encountered or catheter stretches, STOP. Continued pulling could break the catheter.
 - Do not cut or forcefully remove the catheter.
 - After removal, check distal end of catheter for black marking to ensure entire catheter was removed.
3. Cover puncture site with appropriate dressing.
4. Discard catheter per standard hospital protocol.

NOTES

- Product is NOT made with natural rubber latex; however, manufacturing facility may contain natural rubber latex.
- Product is NOT made with DEHP as a plasticizer.

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

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