

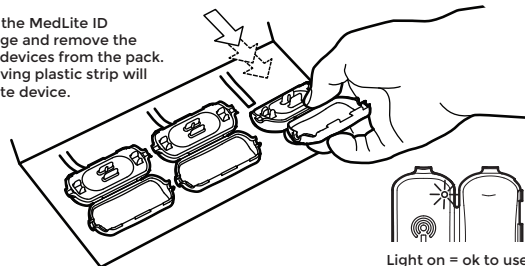
MedLite ID

Primary Medication Line Safety Device

FOR PATIENT SAFETY this product is intended for use with primary medication line only.

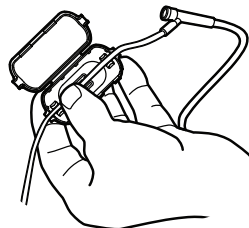
Warning Attach MedLite ID to primary medication line prior to connecting IV to patient.

- 1** Open the MedLite ID package and remove the three devices from the pack. Removing plastic strip will activate device.

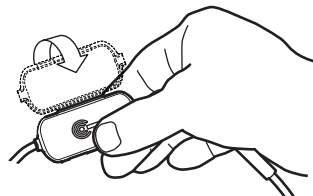


Light on = ok to use

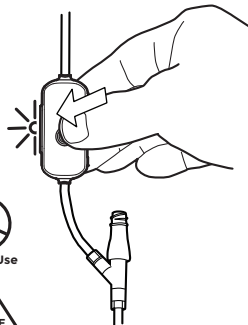
- 2** Prior to connecting the primary medication line to patient, apply MedLite ID devices (see figure) by running the infusion tubing through the tubing grooves on the device



- 3** Inspect tubing for twists or kinks, then close the clamshell door until it clicks.



- 4** Press one of the attached devices to activate the lights for all three devices within a given set.



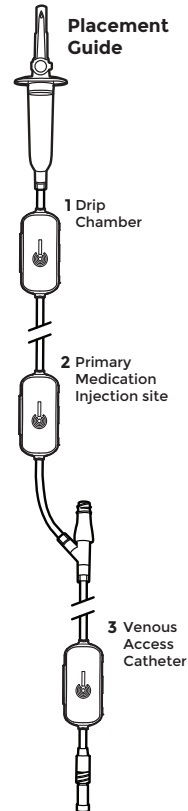
Single Use



Warning
To maintain proper check for pinched or twisted tubing before closing clamshell.

IV bag

Placement Guide



1 Drip Chamber

2 Primary Medication Injection site

3 Venous Access Catheter

Patient



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FOR PATIENT SAFETY this product is intended for use with primary medication line only.

⚠ Warning Attach MedLite ID to primary medication line prior to connecting IV to patient.

⚠ Warning To prevent reduced or obstructed infusion set flow, be sure that when closing the MedLite ID clamshell over the primary medication line that the tubing is not pinched or twisted.

Directions

- 1** Open the MedLite ID peel pack and remove the three devices from the pack.
- 2** Prior to connecting the primary medication line to patient, apply a MedLite ID device below the drip chamber, at the primary medication injection site and near attachment of the venous access catheter (see figure) by running the infusion tubing through the tubing grooves on the device.
- 3** Inspect tubing for twists or kinks, then close the clamshell door until it clicks.
- 4** The MedLite ID device has two forms of identification for the medication line, the attached device itself and an activated light as well.

To activate, press one of the attached devices on the circular squeeze point. This will activate the lights for all three devices within a given set. The devices will turn off after one minute. Pressing on any of the devices will turn all devices back on.

The devices are for one infusion set use and should be disposed of when tubing is changed or treatment completed.

Warning and Disclaimer

This device should be installed by a medical professional. Do not use any component of the device if the product packaging is damaged or opened. MedLite ID is not responsible for use of the product taken from deteriorated packaging, used after expiration of the product shelf life or used on non-medication intravenous lines. MedLite, ID is a single use device with multiple components. Never reuse any portion of the device, even if the device appears undamaged. All portions of the device should be discarded when tubing is changed, or the treatment completed. Care must be taken after opening the device to protect from damage to the components. Prior to installation, all portions of the device must be verified to be in good working order.

If any portion of these instructions are disregarded, the medical professional and their respective employers, agents, members, managers, partners, officers, affiliates, parent entities, assigns, and all other persons or entities who may be responsible for the disregard assume the risk of the use of the product.

By using this product, it is presumed that this warning and disclaimer has been read and understood. For further information or to share comments or complaints about this product, please visit www.medliteid.com.



This device complies with part 15 of the FCC rules. Operation is subject to the following two conditions: (1) This device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation.