






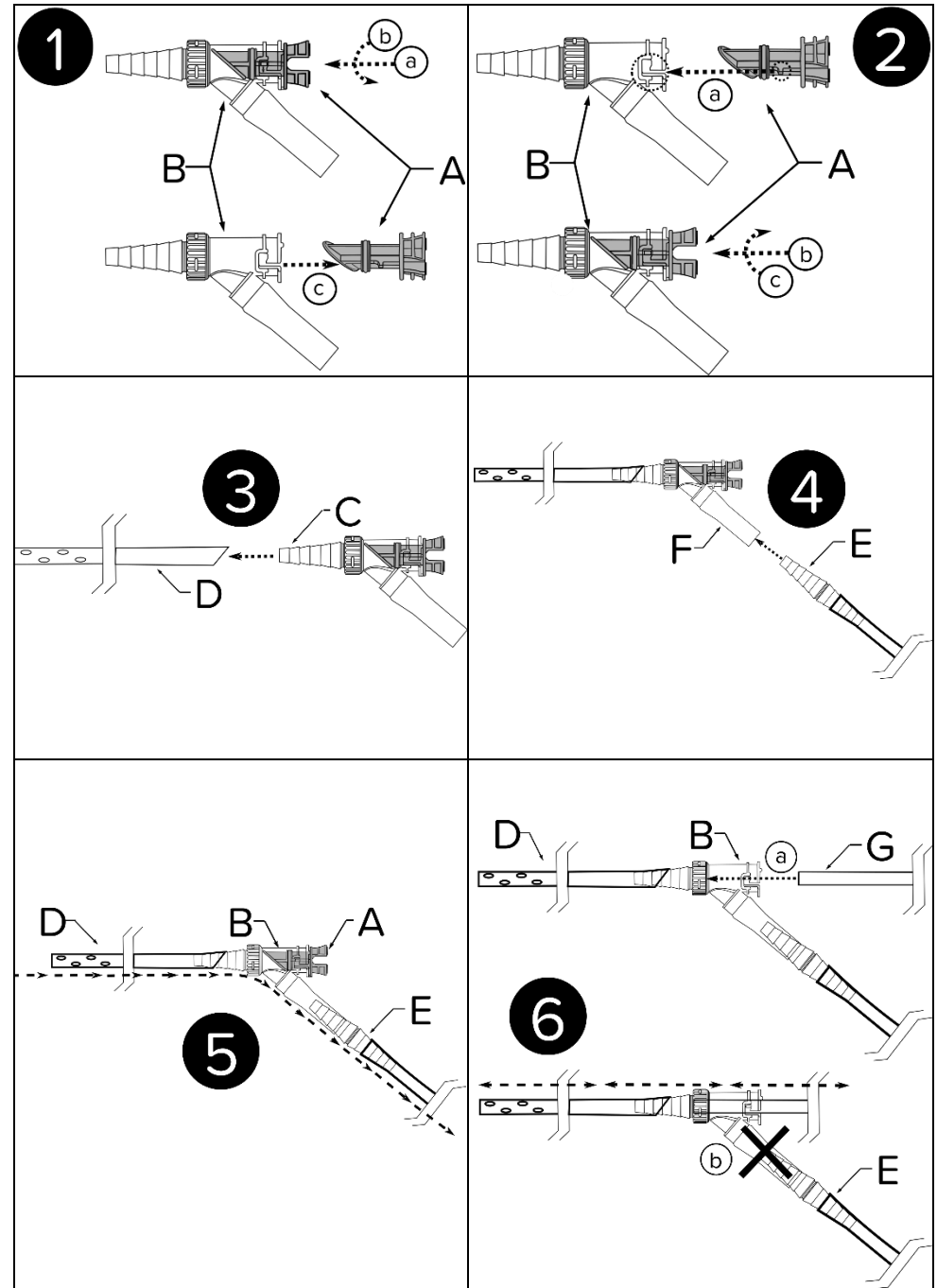
	EN ES FR IT 英语	Not made with natural rubber latex No fabricado con látex de caucho natural. Non fabriqué avec du latex de caoutchouc naturel Non realizzato con lattice di gomma naturale 不是用天然橡膠乳膠製成的
	EN ES FR IT 英语	Do not re-sterilize No volver a esterilizar Ne pas re-stériliser Non ristilizzare 不得重新消毒
	EN ES FR IT 英语	Expiration date Fecha de vencimiento date d'expiration Da utilizzare entro il 有效期
<b>LOT</b>	EN ES FR IT 英语	Lot number Número de lote Numéro de lot Codice partita 批号
<b>DEHP FREE</b>	EN ES FR IT 英语	DEHP free Sin DEHP sans DEHP Privo di DEHP 不含DEHP
	EN ES FR IT 英语	Caution Cuidado Attention Cautela 慎重
	EN ES FR IT 英语	Single patient use only Para uso exclusivo de un sólo paciente À l'usage exclusif d'un seul patient Monouso 限于单个病人使用
<b>STERILE R</b>	EN ES FR IT 英语	Sterilized by irradiation Esterilizado por irradiación Stérilisé par irradiation Sterilizzato con radiazione 放射消毒
<b>Rx Only</b>	EN ES FR IT 英语	Authorized for sale or use by physician only Autorizado para la venta o para el uso sólo por parte de un medico Autorisé à la vente ou à l'usage exclusif d'un médecin Autorizzato solo per vendita o uso da personale medico 医生授权后方可销售或使用
	EN ES FR IT 英语	See instructions for use Consultar las instrucciones de uso Consultez le mode d'emploi Vedere le istruzioni per l'uso 参见使用说明书
	EN ES FR IT 英语	Do not use if package is damaged No usar si el envoltorio está dañado Ne pas utiliser si l'emballage est endommagé Non utilizzare se l'imballaggio è danneggiato 包装损坏后不得使用
	EN ES FR IT 英语	Manufacture date Fecha de fabricación Date de fabrication Data di produzione 生产日期
<u>Manufactured for:</u> CLR® Medical 3901 Calverton Blvd, Suite 155 Calverton, MD 20705 USA TEL: 1-844-758-8270		WWW.CLRMEDICAL.COM

## CLR® Port – Instructions for use



## BEFORE USING THIS PRODUCT, READ THE FOLLOWING INFORMATION THOROUGHLY

### IMPORTANT!

- These instructions are designed to assist with using this product. They should not be used as a guide for the performance of specific surgical techniques.
- This device was designed, tested and manufactured for single use only.
- Reuse of this device may lead to its failure and subsequent patient injury.
- Reprocessing and/or resterilization of this device may create risk of contamination and patient infection and lead to device failure.
- DO NOT reuse, reprocess or resterilize this device.

### DESCRIPTION

The CLR® Port is a Y-shaped catheter adapter configured for connecting to drainage catheters and collection tubing and provides a port for sampling, suction, or infusion.

### INDICATIONS FOR USE

The CLR® Port is intended to provide a sampling, suction, or infusion port for drainage catheters.

### CONTRAINDICATIONS

- This device is not intended for use except as indicated.

### WARNINGS AND PRECAUTIONS

- Procedures should be performed only by medical providers who are trained in surgical techniques and failure modes, precautions, and corrective actions in the event of failure.
- This device should be connected to a regulated suction source. A negative pressure setting that is appropriate for the clinical use should be selected. Similarly, the positive pressure selected should be appropriate for the clinical use. In some cases, the positive pressure will correspond to the height of the column of irrigating fluid.
- This device is provided STERILE and is intended for SINGLE USE ONLY. DISCARD AFTER USE. DO NOT RESTERILIZE.
- Do not use products if the packaging sterile barrier is damaged.
- The CLR® Port is DEHP free, not made with natural rubber latex, and is sterilized using gamma sterilization, except as noted.
- Dispose of all used or damaged products using standard hospital practices for biohazard control.

### INSTRUCTIONS FOR USE

Inspect all components carefully for any damage that may have occurred during shipment, or where applicable, routine handling. Inspect packaging marked as sterile and verify that sterile barriers are intact. Do not use if sterile package is damaged or opened.

*Follow the steps below for use of the CLR® Port, referring to numbered images on Page 1.*

#### Relevant Components

- Access Cap (inserted for normal fluid collection via Collection Tubing, to be removed when requiring access)
- Port Body
- Barbed Catheter Adapter (for connection to Drainage Catheter)
- Drainage Catheter (e.g., chest tube or other drainage catheter)
- Collection Tubing (e.g. tubing from collection unit or suction attachment)
- Tubing Adapter
- Access Catheter (e.g. CLR® Irrigator or other infusion/suction catheter)

#### Access Cap Installation & Removal

- To remove Access Cap:
  - While pushing the Access Cap inward to the Port Body (step **(a)**), twist the Access Cap counter-clockwise so that the tab on the Access Cap aligns with the slot in the Port Body (step **(b)**).
  - Withdraw the Access Cap out of the Port Body (step **(c)**).
- To insert Access Cap:
  - Align the tab on the Access Cap with the slot in the Port Body and insert Access Cap into Port Body (step **(a)**).
  - While pushing the Access Cap inward to the Port Body (step **(b)**), twist the Access Cap clockwise so that the tab on the Access Cap locks into the Port Body (step **(c)**).

#### CLR® Port Connections

- Insert Barbed Catheter Adapter into proximal end of the Drainage Catheter.
- Attach the Collection Tubing to Tubing Adapter.

#### General Usage

- When Access Cap is installed into Port Body, fluids may drain from Drainage Catheter to Collection Tubing.
- When the Access Cap is removed from the Port Body, the Drainage Catheter may be accessed for sampling, suction, or infusion. If a >5 cm long, 30 Fr (10mm) Access Catheter is inserted into Port Body (step **(a)**), the fluid path to the Collection Tubing will be disabled and fluid will only flow between the Drainage Catheter and Access Catheter (step **(b)**).