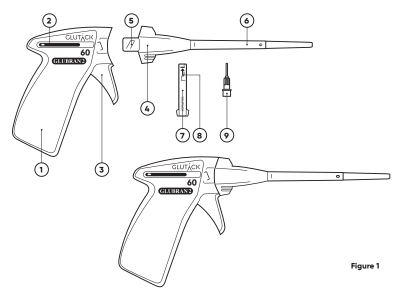
# GLUT<sub>A</sub>CK<sup>®</sup> SHORT

### For the controlled application of Glubran® 2 in open surgery **REF GB-DS SH**



## Features (Description of device)

GLUTACK® SHORT is a Class IIa medical device meeting the requirements of Regulation (EU) 2017/745. The GLUTACK® SHORT device enables the practitioner to apply Glubran® 2 in the form of drops (0.0125 ml - 12.5 mg/ drop) in surgical procedures carried out during open surgery. Mainly designed for the application of Glubran® 2 in the fixation of prostheses during plastic surgery on hernias and incisional hernias.

The sterile, latex-free, single-use device consists of the following components:

- Handpiece (Fig. 1, No. 1) with label stating the number of drops that can be dispensed; equipped with slider (Fig. 1, No.2), which displays the amount of Glubran<sup>®</sup> 2 dispensed during the procedure, and trigger (Fig. 1, No. 3) connected to the gear that forms the drop dispensing system.
- · Catheter consisting of:
- coupling base (Fig. 1, N°4) and chamber for housing the loading cartridge (Fig. 1, N°5);
- long polyethylene shaft, 15 cm (Fig. 1, N° 6) containing a small tube that allows Glubran<sup>®</sup> 2 to pass through; • Cartridge for Glubran® 2 made of transparent plastic (Fig.1,N°7), pre-printed with symbols showing the

direction of insertion into the housing chamber and the loading level (Fig. 1, N°8).

- Transfer tip to be used for loading the cartridge with Glubran<sup>®</sup> 2 (Fig.1,N°9).
   At the end of the directions for assembly, the device is formed by a cannula with handle and trigger
- at the proximal end.

Each individual device can be loaded with the amount of Glubran® 2 that corresponds to the size shown on the package and handle, depending on the various sizes, the amount is therefore enough for at least 30 or 60 drops. It is recommended to use of 0.5 ml Glubran<sup>®</sup>2 with 30 drops of Glutack<sup>®</sup> and the use of 1 ml Glubran<sup>®</sup> 2 with 60 drops Glutack. The green slider, visible from both sides of the handpiece, shows the amount of Glubran® 2 dispensed and the approximate amount remaining. Each time the trigger is pressed/released, one drop (0.0125 ml/drop - 12.5 mg) is delivered from the distal end of the catheter. Read all the directions, precautions and warnings before use. These directions for use only provide information for the correct use of the GLUTACK® SHORT device manufactured by GEM S.r.l, Via dei Campi, 2; 55049 - Viareggio (LU) Italy, for the dispensing drops of Glubran<sup>®</sup> 2.

In the specific case of wall surgery, the device enables the practitioner to apply drops of Glubran<sup>®</sup> 2 on the surface of hernia prostheses.

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In cases involving mesh nets, drops of Glubran® 2 can be dispensed on a prosthesis already adjacent to the trissue, by placing the tip of the GLUACK® SHORT directly into the desired point and gently pressing the trigger. With each click of the trigger, the drop of Glubran® 2 released penetrates the pores of the mesh and begins to polymerise on contact with the moist tissue below.

In cases involving prostheses with film, GLUTACK® SHORT can be used to dispense one or two drops of Glubran<sup>®</sup> 2 directly onto the tissue at the desired point, then bringing the mesh close to the tissue and waiting for complete polymerisation before repeating the application of Glubran® 2 on another area of tissue in the same way.

## Intended use

EN CE 1936

Controlled application of the Glubran® 2 surgical device in the form of drops, during surgical procedures carried out in open surgery

#### **Target population**

Adults

#### Warnings

- ${
  m \Delta}$  The device must only be used by experienced medical practitioners who are suitably trained to use the device.
- ▲ The device is sterile and intended for use on one patient only. Do not reuse, reprocess, clean, disinfect or resterilise the device, as this can lead to a risk of compromising the sterility and performance. ② ③
- ⚠ Make sure that the device has been correctly assembled before use. Assemble according to the instructions in the section on "Directions for assembly and use". [j]
- 🛆 Always check that the system is functioning correctly before use, by dispensing a test drop onto a surgical drape.
- ⚠ The device does not contain user-serviceable parts. Do not try to repair or dismantle the device. If, at any time, the device is found to be damaged or not functioning correctly, discard and replace with another.
- GLUTACK<sup>®</sup> SHORT device is made have been tested with the Glubran<sup>®</sup> 2 medical device only.
- what is shown on this Instruction Sheet.
- ${
  m 
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  m A}$  When applying Glubran $^{\circ}$  2 on the tissue, observe the polymerisation times as shown on the product data sheet.
- ${
  m \Delta}$  During the dispensing process, gently press the trigger and release it at the end of its stroke. Do not continue to increase the pressure on the trigger at the end of its stroke, as this can cause it to break. The break-up.

#### Precautions

- When fixing prostheses in plastic surgery carried out on hernias or incisional hernias, the device must not be used if the prosthetic material is not compatible with cyanoacrylate-based adhesives.
- If the outlet hole of the tip is found to be partly obstructed, clean by rubbing the tip with dry sterile gauze.

## Disposal

Once used, dispose of the device according to local procedures and guidelines.

#### Storage

The device must always be stored in its original packaging. Always store between 5° C (41°F) and 30° C (86°F). \*

## **Expiry date**

The expiry date is shown on the pack.

## Sterility

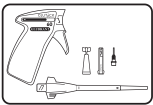
This device is single-use and sterilised with ethylene oxide. STERILE EO

#### Contact

Any serious incident occurring with the device must be reported to the manufacturer GEM srl and to the competent authority of the Member State in which the user and/or patient is established.

## **Product Code**

REF GB- DS SH 30 REF GB- DS SH 60

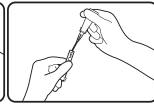


 Arrange the components of the GLUTACK<sup>®</sup> SHORT device (catheter, handpiece, cartridge transfer tip) and a single dose of the Glubran® surgical device on the instrument table

near the neck and applying enough pressure to allow the tip to be inserted. Warning: do not press on the base of the single-dose to avoid leaking of the product.

Open the single dose of Glubran<sup>®</sup> 2. Insert

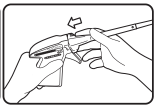
the transfer tip on the single-dose, holding it



Holding the transparent plastic cartridge in a vertical position (black arrow pointing upwards), insert the transfer tip into the small opening. Gradually fill the cartridge by carefully pressing on the body of the single-dose until the product reaches the black line. Once the cartridge is filled, check that it contains no air bubbles.



4) Insert the cartridge into the posterior opening of the catheter (housing chamber) until it reaches the end stop position. When fully inserted, rotate 90 ° clockwise until you hear a click



5) Insert the coupling base (green plastic part with wings) into the handpiece, with the w perpendicular to the handpiece body. ngs





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Directions for assembly and use The device must be adequately prepared and activated to ensure correct functioning.