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CEL+X™ RAPID**Z-Fold Haemostatic Gauze****One Gauze Strip 1.5m x 7.6cm**

Intended Purpose: To be used by trained emergency responders in the pre-hospital setting for temporary treatment of emergency life-threatening bleeding.

Patient Target Group: Adults and children, excluding neonates and infants.

FOR TEMPORARY EXTERNAL USE**STERILE R**

Instructions for
use on back
[www.celoxmedical.com/
instructions-for-use/ifu](http://www.celoxmedical.com/instructions-for-use/ifu)

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(en) Instructions for use:

1 Tear open pack. Before application identify and apply direct pressure on the main part of the bleeding then remove excess blood where practical. Take out the CELOX Rapid and take one end of Z-folded gauze.



2 Tightly pack the unfolding CELOX Rapid directly to the source of the bleeding. Pack remaining wound with CELOX Rapid or standard gauze above skin level.



3 If bleeding persists apply FIRM pressure directly to the wound for 1 minute or until bleeding stops.



4 Excess CELOX Rapid can be torn or cut if necessary. Wrap and tie with a bandage so as to maintain pressure on the wound.

5 Dispose any remaining CELOX Rapid according to local standard protocols for biological waste.

6 Transfer patient to medical facilities as soon as possible.

7 Show empty pack to medical personnel.

ATTENTION MEDICAL FACILITY PERSONNEL:

1. Physically remove CELOX Rapid from the wound and any loose surface granules.
2. Fully flood entire wound area with sterile saline irrigation solution.
3. Proceed with normal irrigation and / or suction.
4. Ensure all product is removed from the wound prior to initiation of wound treatment.
5. Dispose of removed CELOX Rapid and residuals according to local standard protocols for biological waste.

Duration of use: The device and residuals should be removed from the wound within 24 hours from application.

Warnings & Precautions: For external use only. Do not eat. If ingested, drink glass of water to avoid discomfort. Loss of sterility potentially poses a risk of infection. Do not resterilise. Avoid inhalation. Do not apply over eyes. If eye irritation occurs, flush with water for 5 minutes. Keep away from children. Re-use could result in risk of cross-infection and reduced performance. **Contains Chitosan from shellfish – Allergy studies show no adverse reaction. Data on file at Medtrade Products Ltd.**

Contraindications: Do not use in abdominal wounds and wounds unamenable to pressure. Do not pack into body cavities. Device not intended for surgical use.

Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

The Summary of Safety and Clinical Performance (SSCP) for the device is available in the European database on medical devices (Eudamed), where it is linked to the Basic UDI-DI. The Eudamed website is <https://ec.europa.eu/tools/eudamed> and the Basic UDI-DI is 5060206638P000302205U.

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