

bioabsorbable atelocollagen hemostatic felt

Cat. No.:	Size
009	tooth root shape, 10 pieces

LOT

Batch number

REF

Catalog number



Production date



Expiry date



For single use only

STERILE R

Sterilized by gamma-radiation

CE 1023

Notified Body
ITC Zlín, a.s., Czech Republic

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Composition: Bovine atelocollagen type I, 99.9% crystalline

Indication group: Haemostatic

Description: Atelocollagen is a very effective haemostatic. In addition to its excellent properties, Hypro-Sorb Z is well tolerated in the tissue. It is non-immunogenic, resorbable and supports the healing processes.

Activity: Hypro-Sorb Z has a specific activity to thrombocytes and releases clotting factors, which together with plasma binding factors promote fibrinogenesis.

Physiological Effects: Hypro-Sorb Z implanted into the tissue resorbs spontaneously in two to four weeks, depending on the amount and the body region.


Indication: Hypro-Sorb Z is indicated for the control of capillary and parenchymatous bleeding. Hypro-Sorb Z is designed for haemostasis after tooth extraction in stomatology.

Contraindication: Hypro-Sorb Z is not destined for use in instances of pumping arterial haemorrhage. Hypro-Sorb Z must not be left inside the bone fractures fixed by acrylate adhesives because it considerably reduces the bonding strength.

Side Effects: Hypro-Sorb Z is not recommended for persons known to be hypersensitive to the products of bovine origin. Hypro-Sorb Z should not be left between the skin edges as this could interfere with the wound healing and develop wide scars.

In the case of coagulatory disorders or anticoagulation treatment, the haemostatic effect of Hypro-Sorb Z may be reduced or eliminated.

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 **Warning:** Hypro-Sorb Z is sterile. Check original packaging before opening. Do not use if packaging has been opened. Unused Hypro-Sorb Z cannot be re-sterilized. Hypro-Sorb Z is not intended to replace ligation or direct compression in cases of venous bleeding.

Dosage: Hypro-Sorb Z should be used in amounts which fully cover the wounded surface. It is recommended that the surgeon remove the excess felt before closing the wound or leave only the minimum amount in the tissue because it slightly swells and it could exert pressure on the vicinity of the wound. Hypro-Sorb Z implanted into the tissue resorbs spontaneously in two to four weeks depending on its amount and the body region.

Handling instructions: Hypro-Sorb Z is applied usually with its smooth side onto the wound surface, softly pressed down and left on the wound until the fibrine adhesion develops. Bleeding usually stops in two to five minutes; it lasts longer in haemophiliacs. Usually it is used in dry state but it may also be moistened with sterile saline.

Storage conditions: Hypro-Sorb Z must be stored in a dry place and at a normal temperature. Hypro-Sorb Z is not damaged by temperatures from -25°C/-13°F to +50°C/+122°F. It needs to avoid direct sun light.

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Registration: EC Certificate No. 09 0627 QS/NB, EC Design-Examination Certificate No. 09 0628 CN/NB and Declaration of Conformity.

