

Optomesh®

en

INSTRUCTIONS FOR USE

Optomesh®

Surgical hernia mesh

PRODUCT CHARACTERISTICS

Optomesh® non-resorbable surgical mesh products are knitted of transparent and blue monofilament polypropylene yarn using the knitting technique. They are designed for use in abdominal hernia treatment.

The product is offered in various sizes:

-MACROPORE – heavyweight mesh with macro pores;

-THINLIGHT – dünnes, leichtes Netz mit Standard-Porengröße;

-THINLIGHT – dünnes, leichtes Netz mit Standard-Porengröße.

Optomesh mesh implants are available in a full range of sizes. The products do not contain substances of allergenic and toxic origin.

The MACROPORE variant has blue orientation lines facilitating the visibility and to positioning of the product in the operating field.

INDICATIONS

Optomesh® surgical meshes are recommended for reconstructive procedures in order to reinforce the hernia wall with soft tissue defect in the case of abdominal hernias i.e.:

• primary hernia;

• incisional hernias;

• inguinal and femoral hernias;

• umbilical hernias;

• hernias with large gates;

Depending on the hernia type and the applied surgical method the right type of surgical mesh can be chosen as for the surgeon's preferences, nature of reconstruction and applicable medical standards.

CONTRAINDICATIONS

Optomesh® should not be used in infected wounds or susceptible to infection and in conditions where it is not possible to maintain strict surgical asepsis. The Optomesh product is also not recommended in the case of the patient's allergogenicity to the product material (polypropylene).

Optomesh® should not be used in children, women planning pregnancy, pregnant women and when the patient has a contraindication to surgery. It is not recommended to use the Optomesh® in direct contact with internal organs due to the possibility of causing adhesions, fistulas or intestinal obstruction. Do not use the product if there is doubt about its sterility (e.g. cracked, cracked / damaged paper-film packaging, product discolouration).

PRECAUTIONS

1. Optomesh® is designed for use by qualified and trained medical personnel.

2. In order to ensure strict aseptic asperitis during the procedure, it is advisable to use sterilized instruments, appropriate equipment, sterilization of instruments, keeping high personal hygiene) and extraordinary attention while preparing the direct intervention site of the surgeon.

3. Optomesh® sheet only used for non-infected wounds, and in conditions where it is not possible to maintain strict asepsis.

4. Avoid direct contact of the mesh with internal organs.

5. Do not implant the Optomesh® surgical mesh after the expiration date.

6. Optomesh® polypropylene surgical meshes are offered for sale only in a sterile version.

7. Product for single use only.

8. Do not use the product if it is damaged, with structural defects or contaminated.

FIXATION

For fixing the Optomesh® mesh the use of surgical sutures, tackers (surgical screw-like tacks), tissue adhesive or staples (surgical staples) is recommended. Absorbable fixing materials are also allowed. The size of the mesh and distance between the fixing places should be adjusted according to the needs of the patient. The nature of reconstruction and mesh placement should be taken into account when placing the mesh. The fasteners that the implant should be placed to closely adhere to the adjacent tissue. In order to prevent the recurrence of hernia, it is recommended to place the fixing elements at least approx. 1 cm from the edge of the product.

Possible complications

The product application may cause the following complications: exudation of blood and serous fluid, suppuration of the wound, hematomas, abscesses, recurrence of hernia, intense pain (formation of fistulas, adhesions, obstruction).

Diagnosed wound infections should be treated in the fastest possible way using available pharmacological methods. In case of severe and chronic infection, the possibility of partial resection of the infected tissue and/or the mesh should be considered. If there is no progression in the course of the infection, the mesh should be removed. The use of surgical tackers and staples can cause chronic pain resulting from nerve compression. These complications can lead to a prolonged treatment process, chronic pain, recurrent, repeated surgical intervention or, in extreme cases, implant removal. It is not recommended to reuse the product that is non-sterile, damaged or re-used can lead to the above-mentioned complications or serious health damage and patient's life threat.

STERILISATION PROCESS

Sterilisation is carried out by means of ethylene oxide (EO) in a validated process.

The product maintains its sterility within a specified validity period, kept only in the original packaging at maintaining appropriate storage conditions.

STORAGE CONDITIONS

Optomesh® should be stored:

• in dark rooms at humidity 25 - 65%;

• at temperature range 15°C - 35°C;

• under normal conditions, without wetting and mechanical damage or chemical contamination;

in the original unit packaging (box and direct Verpackung). Any mechanical damages to the packaging can expose the product to the loss of sterility.

Date of last verification: 05/2024

de

GE BRAUCHSANWEISUNG

Optomesh®

Chirurgisches Netz zur operativen Behandlung von Hernien

KARAKTERISTIK

Optomesh® nicht resorbierbare chirurgische Netzprodukte werden aus transparentem und blauem Monofilament-Polypropylen gewirkt. Sie sind für die Versorgung von Hernien bestimmt. Diese Erzeugnisse ist in zwei Varianten erhältlich:

-MACROPORE – leichte Gewebe mit makroporösen Poren;

-THINLIGHT – dünnes, leichtes Netz mit Standard-Porengröße.

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The MACROPORE variant has blue Orientation lines facilitating the visibility and to positioning of the product in the operating field.

INDIKATIONEN

Die chirurgischen Netze Optomesh® sind für Rekonstruktionseingriffe zur Auffüllung und/oder Verstärkung der mangelnden Weichgewebe bei abdominalen Hernien empfohlen, die mit einem Gewebe defekt mit soft tissue defect in the case of abdominal hernias i.e.:

• primäre Hernien;

• Incisional Hernien;

• inguinale und femorale Hernien;

• umbilikale Hernien;

• Hernien mit großen Gates;

Depending on the hernia type and the applied surgical method the right type of surgical mesh can be chosen as for the surgeon's preferences, nature of reconstruction and applicable medical standards.

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fr

INSTRUCTIONS D'EMPLOI

Optomesh®

Fil de chirurgie pour le traitement opératoire des hernies

CARACTÉRISTIQUES

Optomesh® n'est pas résorbable chirurgique. Les produits sont fabriqués par technique de tricotage de fil de polypropylène monofilament transparent et bleu. Ils sont destinés à la réparation des hernies abdominales.

Le choix est offert en deux versions:

-MACROPORE – léger et facile à manipuler;

-THINLIGHT – une tige fine et légère avec une grande taille.

Les implants chirurgicaux Optomesh® sont disponibles dans une gamme complète de tailles.

Les produits ne contiennent pas de substances d'origine allergénique et toxique.

La variante MACROPORE offre des lignes d'orientation bleues qui facilitent la visibilité et le positionnement du produit dans le champ opératoire.

INDICATIONS

Les mailles chirurgicales Optomesh® sont conseillées pour les procédures de reconstruction ou de renforcement des défauts des tissus molles en cas de hernie abdominale, d.h.:

• primaire Hernie;

• Hernie inguinale et femorale;

• Hernie umbilicale;

• Hernie avec grands trous;

• Hernie avec élargissement de la ligne blanche;

• Hernie avec grandes enveloppes.

En fonction du type de hernie et de la méthode chirurgicale utilisée, une tige de maille chirurgicale appropriée peut être choisie, compte tenu des préférences du chirurgien, de la nature de la reconstruction et des normes médicales applicables.

AVERTISSEMENTS

Les avis d'interdiction concernant l'utilisation de cette tige chirurgicale doivent être utilisés dans les cas suivants:

- si l'implant n'est pas recommandé pour le patient en raison de son allergie à la matière première (polypropylène);

- si l'implant n'est pas recommandé pour le patient en raison de sa sensibilité à la tige chirurgicale;

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