STERIBON™

Coated, Braided Polyester Suture

DESCRIPTION

STERIBON is sterile, non-absorbable surgical suture made from fine filaments of pure polyester braided, coated to produce a tight smooth uniform strand. STERIBON is dyed with D & C Green number 6 Cl61565 and Coated with Nusil Med 2174 Silicone. Product manufactured in compliance of

STERIBON is crimped with needle (AISI 420 & AISI 300) having the shape and size as 1/2, 3/8, 1/4, 5/8 & Straight.

STERIBON suture is indicated for use in general soft tissue approximation and/or ligation, including in the fields of application where it is surgical practice to employ Nonabsorbable suture.

MODE OF ACTION

STERIBON causes a minimal, acute, inflammatory reaction in tissues, followed by gradual encapsulation of the suture material by fibrous connective tissue.

CONTRAINDICATIONS

Contraindicated in cardiovascular, ophthalmic and neurological procedures.

Device is not clinically evaluated for the using in cardiovascular, ophthalmic and neurological procedures.

When working with Suture materials great care must be taken to ensure that the use of Surgical instruments such as tweezers or needle holders, does not lead to damage by pinching or kinking. The user of Steribon suture materials must be familiar with surgical suture techniques.

APPLICATION

1

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Employ according to surgical requirements.

WARNINGS

Read Instruction for use:-

Read Instruction for use:Use of device is restricted to qualified doctor or paramedic.
Keep out of reach of children. After use of products must be
disposed off as per country law of bio-waste handling rule.
The Firm does not hold any responsibility, if device reused
that may cause serious problems to patients such as HIV,
Hepatitis, contagious Disease/communicable disease etc.
Mishandling or improper transport can cause damage to the
device or packaging. The product is non-toxic and sterile if
the packaging is light? the packaging is intact.

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STERIBON must not be resterilize. Opened, unused packages should be discarded. STERIBON should be stored in between 15°C to 32°C. Do not expose to extreme temperature and humidity. Incase of storage other then above temperature range during any circumstances the intended purpose of device may get effected and this may impact on the performance adsafety.

Do not open the pouch in uncontrolled environment

Do not reuse, may cause infection.

Peters Surgical does not hold any responsibility for any consequences from misuse of the products.

KNOWN CHARACTERISTIC OF DEVICE IN CASE OF RE-USE

Since the cleaning/sterility of the device cannot be guaranteed, chances of infection or disease can transfer

DISPOSAL SYSTEM

Discard the suture in proper waste container & dispose off the product in accordance with accepted medical practice and applicable local, state and country laws and regulations for handling of bio-medical waste

RETURN OF DEVICE

The return of defective device it should be carried out within a week of receipt of the product along with the evidence or damaged product and the product should not have been used under any circumstances, at any condition.

Symbols used on labeling :-

8 **₼** / Mfg.

Do not reuse

Manufacturing year + Month. ☑ / Exp. Use until Year + Month.

LOT STERILE EO

Sterile unless package is opened or damaged. Method of Sterilization-Ethylene Oxide Sterilization.

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Caution Product code.

REF

Consult Instructions for use

Do not resterilize Do not use if package is damaged.



Temperature Limitation.

Authorized European Representative



Moisture Limitation



Keep away from sunlight.



Keep away from moisture. Non-pyrogenic



Single sterile barrier system with protective packaging inside.



Medical device Unique Device Identifier



Patient information website

CE-mark and identification number of notified body. Product conforms to the essential requirements of the Medical Device Directive 93/42/EEC.

TECHNIQUE FOR OPENING OVERWRAP.



Hold pack in left hand, seize foil flap between right hand thumbs and finger, hence exposing transparent plastic flap.



astic flap and paper flap with nands gripping pack between





Now the scrub nurse removes the sterile Primary pouch pack with sterile forceps or with sterile gloved hand.

Recommended technique for ning and handling Primary Foil Pack



Hold sterile primary pouch with the printed top facing towards the scrub nurse. Hold the sterile primary pouch with left hand close to the V-Shaped ($\frac{1}{2}$) cut marked located at the top right corner of the primary pouch.



Tear the pouch from the V-Shaped notch with right hand, down towards the scrub nurse.

Take out the sterile suture winded in suture



PETERS SURGICAL

Immeuble AURÉLIUM 1 Cours de l'Ile Seguin 92100 Boulogne-Billancourt - FRANCE Phone number 00.331.48.10.62.62 Fax number 00.331.48.91.22.99 Email: qualite@peters-surgical.com



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Peters Surgical India (P) Ltd.

169, Sector-4, IMT Manesar, Gurgaon (HR)-122050, INDIA Ph:+91-124-4609500 Fax:+91-124-4609500 (Extn. 137) Email: weera@peters-surgical.com Website: https://peters-surgical.in

