

English**Description**

Prostrolane Blanc_B is a transparent gel supplied in a glass syringe. The product is for single use only. Prostrolane Blanc_B is a sterile medical device, dermal resorbable implant that contains a non-cross-linked Sodium Hyaluronate and peptide complex (Oligopeptide-92, Octapeptide-11, Feruloyl Oligopeptide-33, Valprooyl Oligopeptide-33) Sodium Hyaluronate is non-animal origin and biocompatibility material. The role of Sodium Hyaluronate in the skin is to deliver peptide, to hydrate the skin by holding in water and to act as a cushioning agent. Peptide complex consists of 4 kinds peptides that short chain of amino acids linked by peptide bonds. The role of the peptide complex is to enhance the moisturizing effect, to improve the appearance of photo-aged skin and to delay the degradation of Sodium Hyaluronate. The sodium hyaluronate peptide complex is degradable within 4 weeks. Each carton contains 1 prefilled syringe of 2ml Prostrolane Blanc_B and two traceability labels. (one to be given to the patient, and another to be kept by the doctor in the patient's file)

Composition

PBS/Phosphate Buffered Saline), Sodium Hyaluronate (0.7%), Peptide Complex (Oligopeptide-92, Octapeptide-11, Feruloyl Oligo peptide-33, Valprooyl Oligopeptide-33)

Intended purpose

Prostrolane Blanc_B is indicated for dermal implantation or the treatment of damaged subcutaneous tissue and wrinkles caused by chemical, physical and traumatic effects, as well as sun damaged tissue.

- Site of Application: Face, neck

- Recommended usage volume : 2ml / area (Maximum: 4ml / area)

- Frequency : 1 session / 2 weeks (1 protocol = 2-4 sessions average) (repeat protocol as necessary)

Treatment procedure

Pretreatment Guidelines
Prior to treatment, the patient should avoid taking aspirin, nonsteroidal anti-inflammatory medications, St. John's Wort, or high doses of Vitamin E supplements. These agents may increase bruising and bleeding at the treatment site.

1. It is necessary to counsel the patient and discuss the appropriate indication, risks, benefits and expected responses to the Prostrolane Blanc_B treatment. Advise the patient of the necessary precautions before commencing the procedure.

2. Assess the patient's need for appropriate anesthetic treatment for managing comfort, i.e., topical anesthetic, local or nerve block.

3. The patient's face should be washed with soap and water and dried with a clean towel.

Cleanse the area to be treated with alcohol or another suitable antiseptic solution.

4. Sterile gloves are recommended while using Prostrolane Blanc_B.

5. Before treatment, press rod carefully until a small droplet is visible at the tip.

6. Prostrolane Blanc_B is administered using a thin gauge needle (27G x 13mm or 30G x 13mm) that is sterilized. The sterilized needle is inserted at an approximate angle of 30° parallel to the length of the injected site. Prostrolane Blanc_B should be inserted to the mid to deep dermis. If Prostrolane Blanc_B is injected too superficially this may result in visible lumps and/or bluish discoloration.

7. Inject Prostrolane Blanc_B applying even pressure on the plunger rod. It is important that the injection is stopped just before the needle is pulled out of the skin to prevent material from leaking out or ending up too superficially in the skin.

8. Only correct to 100% of the desired volume effect. Do not overcorrect. If cutaneous deformities the best results are obtained if the defect can be manually stretched to the point where it is eliminated. The degree and duration of the correction depend on the character of the defect treated.

9. Typical usage for each treatment session is specific to the site as well as wrinkle severity. Or to apply on the cleared skin surface, to wait for full absorption.

10. After treatment, stick the traceability label on the patient's medical record.

Direction for Assembly
Assembly by 27G or 30G needle(hypodermic needle) to syringe.

For safe use of Prostrolane Blanc_B, it is important that the needle is properly assembled. Improper assembly may result in separation of the needle and syringe during injection.

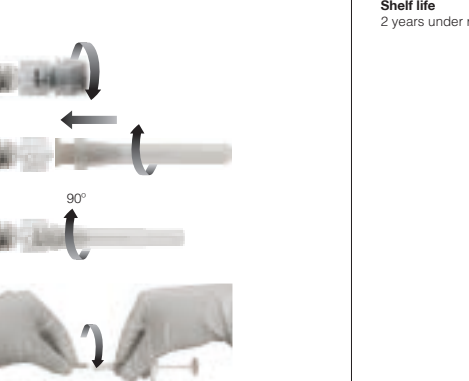
Storage condition
Store between 2°C - 25°C. Protect from freezing and sunlight. Make sure there are no visible signs of damage to the packaging before use.

Shef life
2 years under recommended conditions.

A Manufactured by Caregen Co., Ltd.
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**Spanish****Descripción**

Prostrolane Blanc_B es un gel transparente suministrado en jeringa de cristal. Este producto es de uso único. Prostrolane Blanc_B es un dispositivo médico estéril, dermal resorbible implante que contiene una no-cruzado-cadena de sodio Hialuronato y complejo de péptidos (Oligopeptido-92, Octapeptido-11, Feruloyl Oligopeptido-33, Valprooyl Oligopeptido-33) Sodium Hyaluronate es de origen animal y biocompatibilidad material. El rol de Sodium Hyaluronate en la piel es para entregar el péptido, hidratar la piel en agua y actuar como agente amortiguador. Peptide complex consiste de 4 tipos de péptidos que la cadena corta de aminoácidos unidos por enlaces péptidos. El rol de la complejo de péptidos es para mejorar la hidratación, para mejorar la apariencia de la piel envejecida y para retrasar la degradación de Sodium Hyaluronate. El complejo de péptidos es de 4 tipos de péptidos que la cadena corta de aminoácidos unidos por enlaces péptidos. La función del complejo de péptidos es para mejorar la hidratación, mejorar la apariencia de la piel envejecida y para retrasar la degradación de Sodium Hyaluronate. El complejo de péptidos es de 4 tipos de péptidos que la cadena corta de aminoácidos unidos por enlaces péptidos. La función del complejo de péptidos es para mejorar la hidratación, mejorar la apariencia de la piel envejecida y para retrasar la degradación de Sodium Hyaluronate.

Contraindications

Prostrolane Blanc_B must not be used:
- for the combination with peeling, laser treatment or ultrasound;
- if the patient has cutaneous disorder, inflammation or infection at the treatment site or near to this site;
- in the case of patients have a known hypersensitivity to hyaluronic acid, with a history of severe allergy or anaphylactic shock;
- in case of the patients with autoimmune diseases;
- Due to possible interactions with other filling implants, which have not been researched, it is inadvisable to use Prostrolane Blanc_B into sites in the presence of other filling implants;
- in pregnancy, breast-feeding mother, and in children or minors under 18 ages;
- for patients with bleeding disorders, skin color disorders

Composition

PBS(Phosphate Buffered Saline), Sodium Hyaluronate (0.7%), Peptide Complex (Oligopeptide-92, Octapeptide-11, Feruloyl Oligo peptide-33, Valprooyl Oligopeptide-33)

Intended purpose

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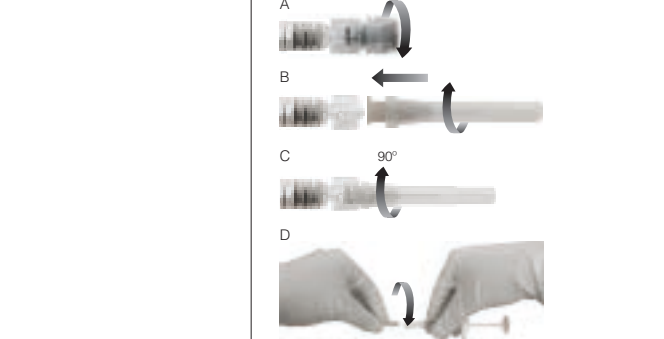
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Storage condition
Store between 2°C - 25°C. Protect from freezing and sunlight. Make sure there are no visible signs of damage to the packaging before use.

Shef life
2 years under recommended conditions.

**Italiano****Descrizione**

Prostrolane Blanc_B è un gel trasparente fornito in una jeringa di vetro. Questo prodotto è per uso unico. Prostrolane Blanc_B è un dispositivo medico sterile, impianto resorbibile dermico che contiene Hialuronato Sódico no reticulato e un complesso di peptidi (Oligopeptido-92, Octapeptido-11, Feruloyl Oligopeptido-33, Valprooyl Oligopeptido-33). Il Hialuronato Sódico è un materiale di origine animale e biocompatibile. Il ruolo del Hialuronato Sódico nella pelle è di fornire peptide, idratare la pelle con l'acqua e funzionare come agente di cuscino. Il complesso di peptidi è costituito da 4 tipi di peptidi che hanno una catena corta di aminoacidi uniti da legami peptidici. Il ruolo del complesso di peptidi è di migliorare l'effetto idratante, di migliorare l'apparenza della pelle invecchiata e di ritardare la degradazione del Hialuronato Sódico. Il complesso di peptidi è un materiale di origine animale e biocompatibile. Il ruolo del Hialuronato Sódico nella pelle è di fornire peptide, idratare la pelle con l'acqua e funzionare come agente di cuscino.

Contraindicazioni

Prostrolane Blanc_B non deve essere usato:
- per la combinazione con peeling, trattamento con laser o ultrasuoni;
- se il paziente tiene un trastorno cutaneo, infiammazione o infettione nel sito di trattamento o vicino a questo sito;
- in caso di pazienti che hanno una notoria ipersensibilità all'acido ialuronico, con una storia di allergia grave o choc anafilattico;
- in caso di pazienti con malattie autoimmuni;
- a causa di possibili interazioni con altri impianti di riempimento, che non sono stati studiati, è consigliabile utilizzare Prostrolane Blanc_B in siti in presenza di altri impianti di riempimento;
- in gravidanza, in allattamento e in bambini o in soggetti di età inferiore ai 18 anni;
- in pazienti con disturbi emorragici, con disturbi del colore della pelle.

Composizione

PBS (fisiologico solfato salsicciano), sodio ialuronato (0,7%), complesso peptidico (Oligopeptido-92, Octapeptido-11, Feruloyl Oligo peptide-33, Valprooyl Oligopeptide-33)

Propósito previsto

Prostrolane Blanc_B è indicato per la implantazione dermica o il trattamento del tejido subcutáneo dañado y las arrugas causadas por efectos químicos, físicos y traumáticos, así como el tejido dañado por el sol.

- Área de aplicación: Cara, cuello.

- Volumen recomendado: 2 ml / área (Máximo: 4ml / área)

- Frecuencia : 1 Sesión / 2 semanas (1 Protocolo = 2-4 Sesiones media) (repetir protocolo si fuera necesario)

Procedimiento

Direcciones de pretratamiento
Antes del tratamiento, el paciente debe evitar tomar aspirina, medicamentos antiinflamatorios no esteroideos, hierba de San Juan, o altas dosis de Suplementos de vitamina E. Estos agentes pueden aumentar los hematomas y sangrado en el área de tratamiento.

1. Es necesario aconsejar al paciente y comentar las indicaciones, riesgos, beneficios y respuestas esperadas al tratamiento con Prostrolane Blanc_B. Aconsejar al paciente de las reacciones locales relacionadas con la implantación de este dispositivo resorbible.

• En caso de que la reacción inflamatoria o cualquier otro efecto secundario no desaparezca en una semana, el paciente debe informar al médico de inmediato.

• Se le solicita al médico que informe cualquier efecto secundario relacionado con Prostrolane Blanc_B a Caregen Co., Ltd. a través de la página web, www.caregen.co.kr o por fax al +82-31-452-3869.

• Deferir el uso de Prostrolane Blanc_B en sitios específicos en los que existe un proceso inflamatorio activo (erupciones cutáneas como costras, granos, erupciones o urticaria) o haya una infección presente hasta que el proceso haya sido controlado.

• Prostrolane Blanc_B está empaquetado para uso en un solo paciente. No volver a esterilizar. No lo use si el paquete es dañado o abierto.

• No usar Prostrolane Blanc_B después de la fecha de caducidad impresa en el paquete.

• Prostrolane Blanc_B debe mezclarse con otros productos antes de la implantación del dispositivo.

• No inyectar en los vasos sanguíneos.

• No inyectar en los párpados.

Precauciones

• En caso de riesgo alérgico, se debe realizar una prueba antes de la primera inyección. Los resultados de la prueba deben adjuntar al historial clínico del paciente.

• El médico debe informar al paciente de las posibles reacciones locales a la jeringa o al dispositivo resorbible.

• Si una reacción inflamatoria o cualquiera otro efecto secundario no desaparece en una semana, el paciente debe informar al médico de inmediato.

• El médico es tenuto a señalar cualquier efecto secundario relevante correlativo a Prostrolane Blanc_B a Caregen Co., Ltd. a través de la Web www.caregen.co.kr o/o via fax al número +82-31-452-3869.

• Revisar la necesidad de consultar al paciente y comentar las indicaciones, riesgos, beneficios y respuestas esperadas al tratamiento con Prostrolane Blanc_B. Informar al paciente de las reacciones locales a la jeringa o al dispositivo resorbible.

• Se necesita aclarar la necesidad de informar al paciente sobre la inyección de Prostrolane Blanc_B.

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