

المحتوى:
قارورة بحجم 10 مل.

التخزين:

ينبغي إغلاق القارورة بعد الاستخدام. يمكن استخدام المنتج المتبقى لمدة تصل إلى 28 يوماً بعد الفتح الأول.

تاريخ انتهاء الصلاحية:

عندما لا تكون العلبة تالفة، لا يجوز الاستخدام بعد تاريخ انتهاء الصلاحية الموضح على العلبة والقارورة. لا يجوز فصل القارورة عن العلبة ونشرة المستهلك، وذلك للحفاظ على كافة المعلومات والتلقيمات الخاصة بالاستخدام الصحيح في نفس المكان.

المورود:

سيريميد الصناعية S.p.A
شارع تيمبيو ديل سيلو 3/5 - 00144 روما (إيطاليا)

المنتج:

مجموعة إروميد م.ض.
شارع تيمبيو ديل سيلو 3/5 - 00144 روما (إيطاليا)

المستورد وصاحب التسجيل:
شركة لابيدوت ميديكال للاستيراد والتوزيع م.ض.
شارع هاشتا 8، قصرين
خدمة الزبائن: 1700-70-90-33

معتمد من قبل وزارة الصحة، قسم المعدات الطبية، رقم التسجيل: 4350729
جهاز طبي مطابق للتوجيه EEC/93/42 بصيغته المعدلة.

إعادة التدوير ينبعي فحص الأنظمة المحلية

يجب إعادة المنتج عن بعد الاستخدام، لا يجوز رمي المنتج في البيئة كنفايات.

التحديث 03 بتاريخ 06/2021

DROP sept

OPHTHALMIC SOLUTION WITH PHLOGOLITHIC ACTION

DROPsept® is an ophthalmic solution, adjuvant in protecting and repairing corneal and conjunctival epithelium, with phlogolithic action. It is indicated for adults and children for the treatment of symptoms such as burning, itching, photophobia, foreign body sensation and redness of the eye, associated with ongoing inflammatory states of the ocular surface and its annexes, such as conjunctivitis, keratitis, blepharitis, dacryocystitis, meibomitis, discomfort and dry eye syndrome.

DROPsept® can be used with wearable contact lenses.

INDICATIONS:

- DROPsept® is indicated as an adjuvant:
- in pre-surgical prophylaxis
 - in the treatment of conjunctival infections
 - in the treatment of corneal infections
 - in the treatment of infections of the ocular annexes (eyelid margins, tear ducts, eyelashes)
 - in the treatment of conjunctivitis
 - in the treatment of keratitis
 - in the treatment of dacryocystitis

INSTRUCTIONS FOR USE:

1. Open the dispenser and remove the protective cap.
2. Gently press the bottle and instill 1-2 drops.

Close the dispenser after use: the remaining product can be used up to 28 days after first opening.
Can be used with wearable contact lenses.

COMPOSITION:

Vitamin E TPGS, Dibasic sodium phosphate, Sodium phosphate monobasic, Sodium chloride, Chlorhexidine digluconate, Purified water.

WARNINGS AND PRECAUTIONS:

- The product is intended for external ophthalmic use only.
- The product is intended for single patient use.
- Do not use in cases of known intolerance or hypersensitivity to its components.
- During product application, do not touch the eye or any other surface with the dispenser tip.
- Do not use in case of damaged or unclosed package before the first opening.
- In case of problems during use, discontinue treatment and consult a doctor.
- In rare cases, phenomena of corneal microcalcifications have been found in patients with previous corneal pathologies and treated with ophthalmic preparations containing phosphates. In this case, it is recommended to use the product under the supervision of the doctor.
- Do not use after the expiry date indicated on the package.
- Before use, carefully read the instructions found on the leaflet.
- Keep away from children.
- Do not waste into the environment after use.

It is possible to report adverse events to the manufacturer via the email address info@iromedgroup.com By reporting adverse events, you can contribute to the safety of this product. Any severe events should be notified to the manufacturer and the competent authority of the country in which this product has been used.

CONTENT:

One 10 ml multidose dispenser.

STORAGE:

Close the dispenser after use: the remaining product can be used up to 28 days after first opening.

EXPIRY DATE:

With an undamaged package, do not use after the expiry date indicated on both the box and the dispenser. Do not separate the dispenser from the box and the leaflet, to keep all the information for a correct use in the same place.

IMPORTER AND REGISTRATION HOLDER:

Lapidot Medical Import and Marketing Ltd.,
8 Hashita St., Caesarea
customer service: 1700-70-90-33

Approved by the Ministry of Health, Medical Device Department. Registration No.: 4350729

MANUFACTURER:
IROMED GROUP S.r.l.
Via Tempio del Cielo 3/5
00144 Rome (Italy)

SUPPLIER:
SERVimed Industrial S.p.A.
Via Tempio del Cielo 3/5
00144 Rome (Italy)

CE 0425 MEDICAL DEVICE conforming to Directive 93/42/EEC as amended.

 Check your local regulations

  

Do not waste into the environment after use

Keep away from children

Rev.03 - 06/2021

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