# Rinoclenil \*100 mcg nasal spray suspension

Bottle for 200 actuations

### COMPOSITION

100 ml of suspension contain:

Active ingredient: beclomethasone dipropionate 77 mg

Excipients: polysorbate 20, microcrystalline cellulose and sodium carboxymethyl cellulose, benzalkonium chloride, phenylethyl alcohol, glucose (dextrose) monohydrate, purified

Each puff delivers 100 micrograms of beclomethasone dipropionate.

# PHARMACEUTICAL FORM AND CONTENT

Nasal spray, suspension, bottle with metering pump and nasal applicator (200 deliveries).

THERAPEUTIC CLASS
Glucocorticoid with nasal decongestive activity for topical use.

## THERAPEUTIC INDICATIONS

Prophylaxis and treatment of seasonal and perennial allergic rhinitis and vasomotor rhinitis.

### CONTRAINDICATIONS

Local viral and tubercular infections. Hypersensitivity to the active ingredient or to any excipient. Contraindicated in children aged under 6 years. Generally contraindicated in pregnancy and lactation (see special warnings).

### PRECAUTIONS FOR USE

The use, especially if prolonged, of products for topical use may give rise to sensitization phenomena and exceptionally to systemic side effects typical of this drug class. In any case, it is necessary to discontinue the treatment and institute a suitable therapy. In pediatric patients receiving prolonged treatments with nasal corticosteroids, a periodical check of their regular growth is recommended.

Though RINOCLENIL® is able to control most cases of seasonal allergic rhinitis, an

abnormally high allergen stimulation can require a proper supplemental therapy, especially

for controlling eye symptoms.
It is necessary to take care of the patients' passage from a systemic steroidal therapy to RINOCLENIL®, if there is any reason to suspect that patients' adrenal function is impaired.

### INTERACTIONS

None known

### SPECIAL WARNINGS

Pregnancy and lactation. In pregnant women the product should be administered in case of real need, under direct medical control. There are insufficient data supporting the safety of use of beclomethasone dipropionate during pregnancy in humans. In reproductive studies in animals, only after high systemic exposures the typical undesired side effects of potent corticosteroids were observed. However, the administration by nasal application of beclomethasone dipropionate avoids the high exposure level occurring with systemic administration. The use of the product during pregnancy should be taken into consideration just when the envisaged benefits outweigh the possible risks for the fetus. The product has been widely used for several years without any apparent damages. It is reasonable to believe that becomethasone dipropionate is excreted in the milk, but with the doses used for nasal application the presence of significant concentrations in the mother milk is unlikely. However, the use of beclomethasone dipropionate during lactation requires that the risk-benefit ratio be duly evaluated both for mother and baby.

METHOD AND FREQUENCY OF ADMINISTRATION Adults and children aged over 6 years: two puffs into each nostril once a day. In children, should it be considered as suitable, an administration scheme of fractioned doses can be maintained, with just one actuation into each nostril twice a day. The onset of action is not immediate, and for a full therapeutic benefit, a regular use is advisable for

The product should not be administered to children aged under 6 years











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Vigorously shake the bottle before each administration. Moreover, before starting therapy, it is recommended to remove the protecting cap (2), protecting ring (3) and repeatedly actuate the metering pump (4) in order to start the nebulization mechanism. Perform the actuation as follows:

1) Carefully clean the nose.

- Remove the protecting cap.

  Remove the side protecting ring blocking the pump.

  Hold the bottle as illustrated in the figure. Repeatedly actuate the metering pump in order to start the nebulization mechanism, to obtain a visible spray.
- Lay the nasal applicator on a nostril, closing with a finger the other one. Breathe in and press at the same time the bottom of the nasal applicator as shown in the figure. In this way a single and exactly metered dose of active ingredient is delivered. Repeat the same operation in the other nostril

  6) After use, close again with the protecting cap and ring. Should the actuator become obstructed, rinse it carefully with lukewarm water, without

intervening on the hole with pointed objects.

The administration of high quantities of beclomethasone dipropionate during a short period of time can bring about suppression of the hypothalamus-hypophysis-adrenal axis. In this case, the dose of RINOCLENIL® should be immediately reduced to the recommended

Systemic undesired effects are extremely unlikely due to the low doses used. Nevertheless systemic undestred effects are extremely unlikely due to the low doses used. Nevertheless, a particular care should be paid in the prolonged use of the product, maintaining the patient under strict control in order to timely detect possible systemic effects (osteoporosis, peptic ulcer, signs of secondary adrenal failure). As for other nasal preparations, some topical burnings might appear, as well as irritation, dryness and seldom epistaxis. Rare cases of nasal septum perforation have been reported following nasal applications with corticosteroids. Seldom cases of intraocular pressure increases or glaucoma have been associated to beclomethasone dipropionate formulations for nasal applications In case of infection, turn to the doctor to allow a suitable therapy to be instituted. The observance of the instructions reported in the present package insert reduces the risk of side effects. Inform your doctor or chemist about any undesired effect even if not reported in the present leaflet

# **EXPIRY DATE AND STORAGE**

See the expiry date printed on the package; this date should be intended for the unopened and correctly stored product.

ATTENTION: do not use the medicinal product after the expiry date indicated on the package.

Keep out of the reach of children

Approved: 4 April, 2003

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ئىكلومىتھازون ۋائى پروپيونىيە 100 مائىكروگرام

100 ملی لیز مسینفن 77 ملی گرام بیکلومیتهازون ذائی پروپیونیٹ کے عال جزیر مشتمل ہے۔ووسرے اجزاء یولی سوربیٹ 20ء مائيكر وكرسفائن سلولوزاورسوۋىم كاربوكسي ميھائل سلولوز ، بينزالكونيم كلورائيۋ، فينائل اسھائل الكومل\_ گلوكوز ( ۋېكىشروز ) موتو مائيڈريٹ ،اور پوريغائيڈ واٹر مِشمَل ہيں۔

رائز کلینل 100 مستقل یامومی الرجیک رینائٹس اورویز دموٹر رینائٹس کی روک تھام اورعلاج کے لئے استعمال کیاجا تا ہے۔

ممانعت برائے علاج:

لوگل دائر کیا ٹیو پر کیلر تھیکھن ۔ را موکلینل 100 ایسے مریضوں کو بھی استعمال نہیں کرنا جائے جن کور بھومیتھا زون ڈائی پروپیونیٹ یا اس میں موجود کسی دوسرے ابڑا ہے و ودحی (Hypersensitivity) ہو۔ تیوسال ہے کم تمریجیاں میں بھی ممانعت ہے۔ حاملہ اور دوده يلانے والي خواتين كوائتيائي گراني اورشد بيضرورت كے تحت استعال كروائيس...

خوراك اورطريقه استعال:

رائز کلینل 100 صرف بڑوں اور 6 سال ہے زیادہ عمر کے بچوں میں استعمال کروا کئیں۔ ہر نتھنے میں دویف (سیرے) دن میں ایک مرتباستعال کریں۔ بچوں میں ایک بنی (سرے) ہر نقینے میں دن میں دومرتباستعال کروائیں۔

تركيب استعال:

استعال ہے میلے ہوتل کواچھی طرح ملائیں۔ 1-ناك كواحتياط المحي طرح صاف كرس - (تصوير 1 ويحييس)

2\_ بوتل کے ڈھکن کواویر ہے ہٹا ئیں۔ (تصویر 2 دیکھیں)

3- يب كوروك والع حطي (Ring) كوبنا كس - (تصوير 3 ديكميس)

4۔ شباوت اور بزی انگلی کونوزل کے دونوں اطراف کالر بر تھیں اور انگو شے کو بوتل کے بنچے تھیں۔ پیپ کو 1-2 دفعہ سپرے کریں تا كه باريك پيوارين جائے۔(تصور 4 ديكھيں)

5۔ ناک کے ایک نتینے میں نوزل لیں اور دوسرے نتینے کوانگی ہے بند کر دیں سیرے کو دیاتے ہوئے کھلے نتینے کے ذریعے سانس اندر تحینچیں۔ایمل کودوس نتھنے میں دوہرائیں۔(تصویر 5 دیکھیں)

6۔استعمال کے بعد عنافتی چھلا (Ring) لگالیں اور نوزل کوؤھکن سے بند کرلیں۔ (نصور 6 دیکھیں)













خوراک ڈاکٹر کی ہدایت کےمطابق استعال کریں۔ صرف متندة اكثر كانسفه كے مطابق عى دوافروفت كى جائے۔ تمام ادومات بجول کی تنا ہے دورر تھیں۔ دواکو 30°C ہے کم ورجہ حرارت برنی اور روشنی سے محفوظ رکھیں۔

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