

TRAVOCORT[®] CREAM

Difflocortolone valerate/Isoconazole nitrate (1mg/10mg)

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Keep this Leaflet. You may need to read it again.

What Travocort[®] Cream is used for

- This medicine is used to treat fungal infections of the skin where inflammation (redness, swelling, soreness) is also a problem.

How Travocort[®] Cream works

- This medicine contains two active substances, isoconazole nitrate and diflucortolone valerate. Isoconazole nitrate treats fungal diseases of the skin and diflucortolone valerate suppresses inflammation of the skin and soothes complaints such as itching, burning and pain.

Before you use Travocort[®] Cream

- When you must not use it

Do not use Travocort[®] Cream if you:

- are allergic (hypersensitive) to isoconazole nitrate or diflucortolone-valerate or any of the other ingredients of this medicine (listed in section 8),
- have skin lesions in the area to be treated which are associated with a tuberculosis or syphilis infection,
- have a viral infection e.g. herpes, shingles or chicken-pox (varicella, herpes zoster),
- have a chronic skin inflammation of the face (rosacea), skin inflammation around the mouth (perioral dermatitis) or a skin reaction following vaccination in the area to be treated.

- Before you start to use it

- Talk to your doctor, pharmacist or nurse before using Travocort.

- Taking other medicine

- Please tell your doctor or pharmacist if you are taking or have recently taken any other medicines, including medicines obtained without a prescription.
- Interactions of Travocort with other medicines are not known so far.

How to use Travocort[®] Cream

- How much to use

- Always use Travocort exactly as your doctor or pharmacist has told you. You should check with them if you are not sure. Do not use Travocort for more than 2 weeks.
- Always wash your hands before and after applying Travocort.

- When to use it

- Travocort should be applied twice daily to the diseased areas of skin. Stop using Travocort when the skin conditions have improved.

- How long to use it

- In general, the duration of treatment must not exceed 2 weeks. If necessary, your doctor may then prescribe a follow-up treatment with a glucocorticoid-free anti-fungal preparation. This applies in particular if Travocort is applied to the groin or the genital area.

- If you forget to use it

- Do not use a double dose to make up for a forgotten dose. When you remember, use the next dose and continue with the treatment as prescribed. See your doctor or pharmacist, if you are worried.

- If you use too much (overdose)

- If you apply too much Travocort once, or accidentally swallow Travocort this is unlikely to be dangerous but contact your doctor or pharmacist if you are worried.

If you have any further questions on the use of this medicine, ask your doctor, pharmacist or nurse.

While you are using it

When using Travocort it is important to know the following:

If you also have a bacterial infection of the skin, your doctor will prescribe you another medicine to use in addition to Travocort, to treat this infection.

- Things you must do

- Regular hygienic measures are essential for successful Travocort treatment: To avoid renewed infection, you should
 - change your personal linen (face cloths, underwear etc. preferably of cotton) daily and boil them,
 - dry the area between the toes thoroughly after washing,
 - change your socks and stockings daily.

If symptoms do not improve, consult the doctor again.

- Things you must not do

- Do not allow Travocort to come into contact with the eyes when applying it to your face. Extensive application of glucocorticoid-containing topical medicines to large areas of the body or for prolonged periods of time, in particular under occlusion (e.g. diapers, dressings) increases the risk of side effects.

Contact your doctor if you experience blurred vision or other visual disturbances.

- Things to be careful of

- There is a risk of developing an eye condition called glaucoma if you apply Travocort covered by a dressing, in large amounts, over a long period of time or if Travocort is applied to the area around the eyes.
- If Travocort is applied to the genital areas, some of its ingredients may cause damage to latex products such as condoms or diaphragms. Therefore, these

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may no longer be effective as contraception or as protection against sexually transmitted diseases such as HIV infection. Talk to your doctor or pharmacist, if you require more information.

Pregnancy and breast-feeding

- If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before using this medicine. Your doctor will carefully weigh the benefits against the risks connected with the use of Travocort.
- Glucocorticoids should not be applied during the first three months of pregnancy to avoid any risk to the development of the unborn baby.
- If you are pregnant, you should in particular avoid applying Travocort covered by a dressing, to large areas of the body or using the cream for a longer time period.
- It is not known whether the active ingredients of Travocort pass into breast milk. A risk to the suckling child cannot be excluded.
- If you are breast-feeding, you should
 - not apply Travocort to the breasts;
 - avoid applying Travocort covered by a dressing or to large areas of the body;
 - avoid using Travocort for a longer time period.
- There are no data showing that fertility is affected by the use of Travocort.

Driving and using machines

- No effects on ability to drive and use machines have been observed in patients treated with Travocort.

Important information about some of the ingredients of Travocort[®] Cream

- Travocort contains cetostearyl alcohol, which may cause local

skin reactions (e.g. contact dermatitis).

Possible side effects

Like all medicines, Travocort can cause side effects, although not everybody gets them.

The following side effects occurred in clinical studies, they are listed according to their frequency:

Common: may affect up to 1 in 10 people

- skin irritation or burning feeling at the application site

Uncommon: may affect up to 1 in 100 people

- redness or dryness at the application site
- stretch marks

Not known: frequency cannot be estimated from the available data

- itching or blisters at the application site
- blurred vision

As with other glucocorticoids that are applied to the skin, such as Travocort, the following local side effects may also occur (their frequency cannot be estimated from the available data):

- Thinning of the skin
- Inflammation of hair follicles
- Increased body hair growth
- Expansion of small superficial blood vessels in the skin
- Skin inflammation around the mouth
- Changes in skin color
- Acne, and/or allergic skin reactions to any of the ingredients of Travocort.

Since the ingredients of Travocort are absorbed by the body through the skin, further side effects to other parts of the body (systemic effects) may occur.

Side effects cannot be excluded in newborns whose mothers have been treated extensively or for a prolonged period of time during pregnancy or while breast-feeding. For example, the activity

of the baby's adrenal glands may be reduced (reduced adrenocortical function) and so the baby's resistance to disease may be lowered.

If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet.

You may report any side effects or adverse drug reactions directly to the National Centre for Adverse Drug Reaction Monitoring by visiting the website npra.gov.my [Consumers→Reporting Side Effects to Medicines (ConSERF) or Vaccines (AEFI)]

By reporting side effects you can help provide more information on the safety of this medicine.

Storage and Disposal of Travocort[®] Cream

- Storage

Store Travocort out of the sight and reach of children.

Do not store above 30°C.

Do not use this medicine after the expiry date which is stated on the tube and carton after 'EXP'. The expiry date refers to the last day of that month.

- Disposal

Do not throw away any medicine via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

Product Description

- What it looks like

Travocort[®] Cream is a white to yellowish opaque cream.

- Ingredients

- Active ingredient:
Diflucortolone valerate and isoconazole nitrate. Travocort

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contains 0.1% w/w diflucortolone valerate (1mg/g) and 1% w/w isoconazole nitrate (10mg/g).

- Inactive ingredient:

The other ingredients are:

- Paraffin, white soft
- Paraffin, liquid
- Cetostearyl alcohol
- Polysorbate 60
- Sorbitan stearate
- Disodium edetate dihydrate
- Water, purified

See end of Section 5 for cetostearyl alcohol advice.

- MAL number(s):

Travocort[®] Cream:
MAL20001796AZ

Manufacturer

LEO Pharma Manufacturing Italy S.r.l.
Via E. Schering 21
20054 Segrate (Milan) Italy

Product Registration Holder

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