הודעה על החמרה (מידע בטיחות) בעלון לרופא

השינויים בעלון <mark>מסומנים על רקע צהוב</mark>

פרטים על השינוי/ים המבוקש/ים		
טקסט חדש	טקסט נוכחי	פרק בעלון
Protopic 0.03% ointment is indicated in adults, adolescents and children from the age of 2 years. Flare treatment Adults and adolescents (16 years of age and above) Treatment of moderate to severe atopic dermatitis in adults who are not adequately responsive to or are intolerant of conventional therapies such as topical corticosteroids. Children (2 years of age and above) Treatment of moderate to severe atopic dermatitis in children (2 years of age and above) who failed to respond adequately to conventional therapies such as topical corticosteroids. Maintenance treatment Treatment of moderate to severe atopic dermatitis for the prevention of flares and the prolongation of flare-free intervals in patients experiencing a high frequency of disease exacerbations (i.e. occurring 4 or more times per year) who have had an initial response to a maximum of 6 weeks treatment of twice daily tacrolimus ointment (lesions cleared, almost cleared or mildly affected).	Treatment of moderate to severe atopic dermatitis in adults who are not adequately responsive to or are intolerant of conventional therapies such as topical corticosteroids. Treatment of moderate to severe atopic dermatitis in children (2 years of age and above) who failed to respond adequately to conventional therapies such as topical corticosteroids.	Indications
The effect of treatment with Protopic ointment on the developing immune system of children aged below 2 years, especially the young, has not yet been established and this should be taken into account when prescribing to this age group (see section 4.1). The use of tacrolimus ointment is not recommended in patients with a skin barrier defect, such as Netherton's syndrome, lamellar ichthyosis, generalized erythroderma or cutaneous Graft Versus Host Disease. These skin conditions may increase systemic absorption of tacrolimus. Oral use of acrolimus is also not recommended to treat these skin conditions. Post-marketing cases of increased acrolimus blood level have been reported in these conditions. The use of Protopic ointment in patients with genetic epidermal barrier defects such as Netherton's	The effect of treatment with Protopic ointment on the developing immune system of children, especially the young, has not yet been established and this should be taken into account when prescribing to this age group (see section 4.1). The use of Protopic ointment in patients with genetic epidermal barrier defects such as Netherton's syndrome is not recommended due to the potential for permanently increased systemic absorption of tacrolimus. The safety of Protopic ointment has not been established in patients with generalised erythroderma.	Warnings and Precautions

syndrome is not recommended due to the potential

for permanently increased systemic absorption of tacrolimus. The safety of Protopic ointment has not been established in patients with generalised erythroderma.		Ocutario dicatione
Paediatric population An interaction study with protein-conjugated vaccine against Neisseria menigitidis serogroup C has been investigated in children aged 2-11 years. No effect on immediate response to vaccination, the generation of immune memory, or humoral and cell-mediated immunity has been observed (see section 5.1).		Pediatrics
There are no fertility data available. Pregnancy There are no adequate data from the use of tacrolimus ointment in pregnant women. Studies in animals have shown reproductive toxicity following systemic administration (see section 5.3). The potential risk for humans is unknown. Protopic ointment should not be used during pregnancy unless clearly necessary. Breastfeeding Human data demonstrate that, after systemic administration, tacrolimus is excreted into breast milk. Although clinical data have shown that systemic exposure from application of tacrolimus ointment is low, breast-feeding during treatment with Protopic ointment is not recommended. There are no adequate data from the use of tacrolimus ointment in pregnant women. Studies in animals have shown reproductive toxicity following systemic administration (see section 5.3). The potential risk for humans is unknown. Protopic ointment should not be used during pregnancy unless clearly necessary. Human data demonstrate that, after systemic administration, tacrolimus is excreted into breast milk. Although clinical data have shown that systemic exposure from application of tacrolimus ointment is low, breast feeding during treatment with Protopic ointment is not recommended.	There are no adequate data from the use of tacrolimus ointment in pregnant women. Studies in animals have shown reproductive toxicity following systemic administration (see section 5.3). The potential risk for humans is unknown. Protopic ointment should not be used during pregnancy unless clearly necessary. Human data demonstrate that, after systemic administration, tacrolimus is excreted into breast milk. Although clinical data have shown that systemic exposure from application of tacrolimus ointment is low, breast-feeding during treatment with Protopic ointment is not recommended.	Fertility, Pregnancy and Lactation
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	General disorders and administration site conditions Very common: Application site burning, application site pruritus Common: Application site warmth, application site erythema, application site pain, application site irritation, application site paraesthesia, application site rash	Adverse events

aetiolo	Infections and infestations
gy includi ng but not limited to: Eczem a herpeti cum,	Common: Herpes viral infections (herpes simplex dermatitis [eczema herpeticum], herpes simplex [cold sores], Kaposi's varicelliform eruption)
Follicu litis, Herpes simple x, Herpes virus infecti on, Kaposi 's varicell iform	Skin and subcutaneous tissue disorders Common: Folliculitis, pruritus Uncommon: Acne Nervous system disorders Common: Paraesthesias and dysaesthesias (hyperaesthesia, burning sensation) Metabolism and nutrition disorders
Metab olism and intoler ance nutritio ance (facial disorde flushin rs g or skin irritatio n after consu mption of an alcohol ic bevera ge)	Common: Alcohol intolerance (facial flushing or skin irritation after consumption of an alcoholic beverage) The following adverse reactions have been reported during post- marketing experience: Skin and subcutaneous tissue disorders: Rosacea.
Nervou s hesias system and disorde dysaest rs hesias (hyper aesthes ia, burnin g sensati on) Skin Pruritu Acne*	Rosacea*
and subcut aneous tissue disorde rs	Application site oedema*

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	rash	
Investi	Drug level	
gations	increased*	
gations	(see section	
	4.4)	
	reaction has been reported during post-	
marketing expe	erience	
<u>General disord</u>	ers and administration site conditions	
Very common:	: Application site burning,	
	application site pruritus	
Common:	Application site warmth.	
ommon.	application site erythema	
	application site or thema;	
	application site purning, application site pruritus Application site warmth, application site erythema, application site pain, application site irritation, application site paraesthesia, application site	
	she irritation, application site	
	paraesthēsia, application site	
	rash	
infections and	infestations	
Common:	Herpes viral infections (herpes	
	simplex dermatitis [eczema	
	herpeticum], herpes simplex	
	simplex dermatitis [eczema herpeticum], herpes simplex [cold sores], Kaposi's varicelliform eruption)	
	varicelliform eruption)	
Skin and subcu	<mark>itaneous tissue disorders</mark>	
Common:	Folliculitis, pruritus	
Jncommon:		
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T	4:	
Vervous syster	n disorders	
Common:	Paraesthesias and dysaesthesias	
	(hyperaesthesia, burning	
	sensation)	
Metabolism an	d nutrition disorders	
Common:	nd nutrition disorders Alcohol intolerance (facial flushing or skin irritation after consumption of an alcoholic	
	flushing or skin irritation after	
	consumption of an alcoholic	
	beverage)	

beverage)
The following adverse reactions have been reported

during post marketing experience: Skin and subcutaneous tissue disorders: Rosacea. Paediatric population Frequency, type and severity of adverse reactions in children are similar to those reported in adults.		
A potential interaction between vaccination and application of Protopic eintment has not been investigated. Because of the potential risk of vaccination failure, vaccination should be administered prior to commencement of treatment, or during a treatment free interval with a period of 14 days between the last application of Protopic and the vaccination. In case of live attenuated vaccination, this period should be extended to 28 days or the use of alternative vaccines should be considered.	A potential interaction between vaccination and application of Protopic ointment has not been investigated. Because of the potential risk of vaccination failure, vaccination should be administered prior to commencement of treatment, or during a treatment-free interval with a period of 14 days between the last application of Protopic and the vaccination. In case of live attenuated vaccination, this period should be extended to 28 days or the use of alternative vaccines should be considered.	Drug Interactions
Protopic ointment is administered topically and is unlikely to have an effect on the ability to drive or use machines. No studies on the effects on the ability to drive and use machines have been performed. Protopic ointment is administered topically and is unlikely to have an effect on the ability to drive or use machines.	No studies on the effects on the ability to drive and use machines have been performed. Protopic ointment is administered topically and is unlikely to have an effect on the ability to drive or use machines.	Effects on ability to drive and use machines
Protopic treatment should be initiated by physicians with experience in the diagnosis and treatment of atopic dermatitis. Protopic can be used for short term and intermittent long term treatment. Treatment should not be continuous Protopic is available in two strengths, Protopic 0.03% and Protopic 0.1% ointment. Posology Flare treatment Protopic can be used for short-term and intermittent long-term treatment. Treatment should not be continuous on a long-term basis. Protopic treatment should begin at the first appearance of signs and symptoms. Each affected region of the skin should be treated with Protopic until lesions are cleared, almost cleared or mildly affected. Thereafter, patients are considered suitable for maintenance treatment (see below). At the first signs of recurrence (flares) of the disease symptoms, treatment should be re-initiated. Adults and adolescents (16 years of age and above) Treatment should be started with Protopic 0.1% twice a day and treatment should be continued until clearance of the lesion. If symptoms recur, twice daily treatment with Protopic 0.1% should be restarted. An attempt should be made to reduce the frequency of application or to use the lower strength Protopic 0.03% ointment if the clinical condition allows. Generally, improvement is seen within one week of starting treatment. If no signs of improvement are seen after two weeks of treatment, further treatment options should be considered.	Protopic should be initiated by physicians with experience in the diagnosis and treatment of atopic dermatitis. Protopic can be used for short-term and intermittent long-term treatment. Treatment should not be continuous Generally, improvement is seen within one week of starting treatment. If no signs of improvement are seen after two weeks of treatment, further treatment options should be considered. Protopic ointment should be applied as a thin layer to affected or commonly affected areas of the skin. Protopic ointment may be used on any part of the body, including face, neck and flexure areas, except on mucous membranes. Protopic ointment should not be applied under occlusion (see section 4.4). Protopic is not recommended for use in children below the age of 2 years until further data are available. Specific studies have not been conducted in elderly patients. However, the clinical experience available in this patient population has not shown the necessity for any dosage adjustment. Treatment Protopic treatment should begin at the first appearance of signs and symptoms. Each affected region of the skin should be treated with Protopic until lesions are cleared, almost cleared or mildly affected. At the first signs of recurrence (flares) of the disease symptoms, treatment should be reinitiated.	Dosage and Administration

Elderly patients

Specific studies have not been conducted in elderly patients. However, the clinical experience available in this patient population has not shown the necessity for any dosage adjustment.

Paediatric population

Children (2 years of age and above) should use the lower strength Protopic 0.03% ointment. Treatment should be started twice a day for up to three weeks. Afterwards the frequency of application should be reduced to once a day until clearance of the lesion (see section 4.4).

Protopic ointment should not be used in children aged below 2 years until further data are available.

Maintenance treatment

Patients who are responding to up to 6 weeks treatment using tacrolimus ointment twice daily (lesions cleared, almost cleared or mildly affected) are suitable for maintenance treatment.

Adults and adolescents (16 years of age and above)
Adult patients should use Protopic 0.1% ointment.
Protopic ointment should be applied once a day twice weekly (e.g. Monday and Thursday) to areas commonly affected by atopic dermatitis to prevent progression to flares. Between applications there should be 2–3 days without Protopic treatment.

After 12 months treatment, a review of the patient's condition should be conducted by the physician and a decision taken whether to continue maintenance treatment in the absence of safety data for maintenance treatment beyond 12 months.

If signs of a flare reoccur, twice daily treatment should be re-initiated (see flare treatment section above).

Elderly patients

Specific studies have not been conducted in elderly patients (see flare treatment section above).

Paediatric population

Children (2 years of age and above) should use the lower strength Protopic 0.03% ointment.

Protopic ointment should be applied once a day twice weekly (e.g. Monday and Thursday) to areas commonly affected by atopic dermatitis to prevent progression to flares. Between applications there should be 2–3 days without Protopic treatment.

The review of the child's condition after 12 months treatment should include suspension of treatment to assess the need to continue this regimen and to evaluate the course of the disease.

Protopic ointment should not be used in children aged below 2 years until further data are available.

Method of administration

Protopic ointment should be applied as a thin layer to affected or commonly affected areas of the skin. Protopic ointment may be used on any part of the body, including face, neck and flexure areas, except on mucous membranes. Protopic ointment should not be applied under occlusion because this method of administration has not been studied in patients (see section 4.4).

Use in children (2 years of age and above) Treatment should be started twice a day for up to three weeks. Afterwards the frequency of application should be reduced to once a day until clearance of the lesion (see section 4.4).

Use in adults (16 years of age and above) Protopic is available in two strengths, Protopic 0.03% and Protopic 0.1% ointment. Treatment should be started with Protopic 0.1% twice a day and treatment should be continued until clearance of the lesion. If symptoms recur, twice daily treatment with Protopic 0.1% should be restarted. An attempt should be made to reduce the frequency of application or to use the lower strength Protopic 0.03% ointment if the clinical condition allows.

Protopic ointment should be applied as a thin layer to affected or commonly affected areas of the skin. Protopic ointment may be used on any part of the body, including face, neck and flexure areas, except on mucous membranes. Protopic ointment should not be applied under occlusion (see section 4.4). Protopic is not recommended for use in children below the age of 2 years until further data are available. Specific studies have not been conducted in elderly patients. However, the clinical experience available in this patient population has not shown the necessity for any dosage adjustment. **Treatment** Protopic treatment should begin at the first appearance of signs and symptoms. Each affected egion of the skin should be treated with Protopic until lesions are cleared, almost cleared or mildly affected. At the first signs of recurrence (flares) of the disease symptoms, treatment should be reinitiated. Use in children (2 years of age and above) Treatment should be started twice a day for up to three weeks. Afterwards the frequency of application should be reduced to once a day until clearance of the lesion (see section 4.4). Use in adults (16 years of age and above) Protopic is available in two strengths, Protopic 0.03% and Protopic 0.1% ointment. Treatment should be started with Protopic 0.1% twice a day and treatment should be continued until clearance of the lesion. If symptoms recur, twice daily treatment with Protopic 0.1% should be restarted. An attempt should be made to reduce the frequency of application or to use the lower strength Protopic 0.03% ointment if the clinical condition allows. A seven-month, double blind, randomised parallel group study of paediatric patients (2-11 years) with moderate to severe atopic dermatitis was performed. In one arm patients received Protopic 0.03% ointment (n=121) twice a day for 3 weeks and thereafter once a day until clearance. In the comparator arm patients received 1% hydrocortisone acetate ointment (HA) for head and neck and 0.1% hydrocortisone butyrate ointment for trunk and limbs (n=111) twice a day for 2 weeks and subsequently HA twice a day to all affected areas. During this period all patients and control subjects (n=44) **Pharmacodynamics** received a primary immunisation and a rechallenge with a protein-conjugate vaccine against Neisseria *menigitidis* serogroup C. The primary endpoint of this study was the response rate to vaccination, defined as the percentage of patients with a serum bactericidal antibody (SBA) titre ≥ 8 at the week 5 visit. Analysis of the response rate at week 5 showed equivalence between the treatment groups (hydrocortisone 98.3%, tacrolimus ointment 95.4%; 7-11 years: 100% in both arms). The results in the control group were similar.

The primary response to vaccination was not affected.	
Paediatric population The pharmacokinetics of tacrolimus after topical application are similar to those reported in adults, with minimal systemic exposure and no evidence of accumulation (see above).	Pharmacokinetics
	Instructions for use and handling and disposal