LOCORTEN VIOFORM®
Clioquinol/flumetasone pivalate

Qualitative and quantitative composition

A clear yellowish solution with a volume of 7.5ml (32 ± 3 drops per 1ml) containing 1% clioquinol and 0.02% flumetasone pivalate.

Polyethylene glycol 300. Flumetasone is a synthetic difluorinated glucocorticoid. Clioquinol is a halogenated hydroxyquinolone derivative.

Pharmaceutical form

Ear drops

Clinical particulars

Therapeutic indications

Eczema of the external auditory meatus in which secondary infection with microorganisms sensitive to clioquinol has occurred.

- Otitis externa
- Otomycosis.

Dosage and method of administration

Before application, the auditory meatus should be cleansed and dried carefully.

Instil 2 or 3 drops twice daily into the auditory meatus by gently squeezing the plastic bottle. The patient should be either sitting or lying down with the treated ear turned upwards during application. This position should be maintained for at least 1-2 minutes following the application.

Alternatively, a gauze or cotton wick saturated with the solution may be inserted into the ear canal. Keep the wick moistened by adding further solution. It should be replaced at least once every 24 hours.

The solution may be warmed to body temperature prior to each application (e.g. by holding the bottle in the hands). Heating above body temperature should be avoided. Contamination of the dropper with material from the ear, fingers, or other sources should be avoided. Treatment should normally not exceed 10 days.

LOCORTEN-VIOFORM® is dissolved in a polyethylene glycol vehicle which forms an inert, non-irritant, rather viscous medium. This medium has a softening effect on the cerumen and does not prevent drainage, thus ensures prolonged contact of the active ingredients with the surface of the ear canal.

If no improvement occurs within about 1 week, the therapy should be discontinued; it is then advisable to identify the pathogens and to institute an appropriate treatment. Use of the product is not advisable in children under the age of 2 years.

Contraindications

- Perforation of the ear-drum (suspected or verified)
• Application to the eye
• Viral infections of the skin (e.g. varicella, herpes simplex, herpes zoster, common warts, plane warts, condylomata, mollusca contagiosa)
• Syphilitic skin affections, tuberculosis of the skin, bacterial skin infections (e.g. pyoderma)
• Fungal and yeast infections of skin
• Disorders associated with therapy-resistant secondary infections
• Known hypersensitivity to flumetasone pivalate, known hypersensitivity to clioquinol, hydroxyquinolines and other quinoline derivatives, to iodine, as well as to other components contained in LOCORTEN-VIOFORM® ear drops.
• Use in children under 2 years of age.

Special warnings and special precautions for use

Prior to the beginning of therapy, the ear-drum should be checked by the physician. If there is a risk that perforation of the ear-drum may occur or is perforated, LOCORTEN-VIOFORM® ear drops should not be used.

If no improvement occurs within about 1 week, the therapy should be discontinued; it is then advisable to identify the pathogens and to institute an appropriate treatment (see also “Dosage and method of administration”).

LOCORTEN-VIOFORM® ear drops should not be allowed to come into contact with the conjunctiva, due to the possibility of contamination of the conjunctiva with the risk of developing open-angle glaucoma or subcapsular cataract.

Contact with LOCORTEN-VIOFORM® ear drops may cause discolouration of the hair and of clothing and bed linen.

Topical use of clioquinol-containing preparations may lead to a marked increase in protein-bound iodine (PBI) (see also "Overdosage") in patients with normal thyroid function and therefore may interfere with tests of thyroid function (such as PBI, radioactive iodine and butanol extractable iodine). Other thyroid function tests, such as the T3 resin sponge test or T4 determination, are unaffected.

LOCORTEN-VIOFORM® should not be used concomitantly with products containing heavy metals, as clioquinol forms coloured complex-bound salts with these substances.

In the event of suspected allergic skin reaction, LOCORTEN-VIOFORM® should be withdrawn.

Topical corticosteroids are absorbed systemically, and the risk of systemic adverse reactions should be considered, especially with high dosage or long-term treatment. It will be necessary to stop the treatment and start appropriate therapy.

Corticosteroids may mask or exacerbate an existing infection. Long-term use may compromise the immune system, increasing the risk of secondary infections.

It is not advisable to use it continuously for more than five days without consulting the doctor. In cases of superinfection, the product should be withdrawn and appropriate measures should be taken.

Medicines should be kept out of the reach of children. Use of the product is not advisable in children under the age of 2 years (see also “Dosage and method of administration”).

Interaction with other medicaments and other forms of interaction

Topical use of clioquinol-containing preparations may increase the amount of protein-
bound iodine (PBI) in patients with normal thyroid function and therefore may interfere with tests of thyroid function (such as PBI, radioactive iodine and butanol extractable iodine). Such tests should therefore be performed no sooner than one month after cessation of therapy. Other thyroid function tests, such as the T3 resin sponge test or T4 determination, are unaffected.

The ferric chloride test for phenylketonuria may yield a false-positive result when clioquinol is present in the urine.

However, no similar reports with LOCORTEN-VIOFORM® ear drops have been received to date.

**Pregnancy and lactation**

**Pregnancy**
There is limited information on use of LOCORTEN-VIOFORM® ear drops in pregnant women. Animal experiments relevant to the safety assessment of corticosteroids, although not specifically conducted with LOCORTEN-VIOFORM® ear drops, have shown either teratogenic potential or other adverse effects on the embryo and/or the foetus. However, no reports on adverse effects with LOCORTEN-VIOFORM® ear drops in human pregnancy have been received to date.

The safety of use during pregnancy has not been demonstrated. Although systemic exposure to the active ingredients is expected to be low following local application in the ear. When using LOCORTEN-VIOFORM® ear drops in pregnancy, the risk-benefit relationship must be carefully considered.

Corticosteroids are known to cross the placenta and therefore potentially influence the foetus. This will be significant predominantly when intensive treatment is administered to large areas using a potent or highly potent product.

**Lactation**
It is not known whether the active substances of LOCORTEN-VIOFORM® ear drops and/or their metabolite(s) pass into the breast milk when the preparation is applied topically.

For safety reasons caution is indicated.

**Effects on ability to drive and use machines**
None known to date. But Locacorten-Vioform® is not expected to have an effect.

**Adverse effects**

The following adverse drug reactions have been derived from post marketing experience with LOCORTEN-VIOFORM® ear drops. Because these reactions are reported voluntarily from a population of uncertain size, it is not possible to reliably estimate their frequency which is therefore categorized as not known. The adverse drug reactions are listed according to system organ classes in MedDRA. Within each system organ class, ADRs are presented in order of decreasing seriousness.

**Infections and infestations:**
Not known: Manifestations of bacterial infections, viral and fungal infections.

**Immune system disorders:**
Occasional: Hypersensitivity

**Eye disorder:**
Not known: Increased risk of cataract*
Skin and subcutaneous tissue disorders:
Not known: Skin atrophy* (Often irreversible), telangiectasia*, purpura*, skin striae*, rosacea-like dermatitis and dermatitis (perioral) with or without skin atrophy*, dermatitis contact (due to components of the vehicle or due to the corticosteroid itself)*, skin depigmentation*, hypertrichosis*, hair colour changes (contact with Locorten-Vioform® ear drops)

General disorders and administration site conditions:
Occasionally: application site irritation, application site burn, application site pruritus application site rash, rebound effect, which may lead to steroid dependence*, impaired healing*

Treatment should be discontinued if severe irritation or sensitisation develops.

Investigation:
Not known: Intraocular pressure increased*

*The localised undesirable effects that have been reported after topical application of corticosteroids in general.

Side effects observed after prolonged periods of application

Endocrine disorders:
Not known: Adrenal suppression

The risk of localised undesirable effects increases with potency of the product and duration of treatment. Application in skin folds increases this risk. The face, hairy skin and genital skin are especially sensitive to localised effects.

Incorrect use may mask and/or worsen bacterial, parasitic, fungal and viral infections. In addition, the development of resistant organisms requires the initiation of appropriate treatment and, should this fail, cessation of treatment with the product.

Systemic undesirable effects due to topical application of corticosteroid preparations are rare in adults, but may be serious. The risk of systemic effects is the greatest with:
• application under occlusion,
• application in skin folds,
• application on large areas of skin,
• prolonged application,
• application in children (thin skin and relatively large skin surface make children highly sensitive).

Overdose
Treatment with clioquinol-containing preparations applied to extensive or eroded areas of skin may already within 1 week lead to increased PBI values and to signs and symptoms resembling those of thyrotoxicosis. Elevated PBI values also occur where relatively small areas of skin are treated for more than 1 week.

However, no similar reports with LOCORTEN-VIOFORM® ear drops have been received to date.

No cases of acute poisoning have been reported to date. However, the possibility of such an occurrence owing to accidental ingestion by children cannot be excluded. This would result in gastrointestinal disorders accompanied by vomiting and nausea. There are no known specific antidotes.

Pharmacological properties
Pharmacodynamic properties

Pharmacotherapeutic group: Otologicals corticosteroids and anti infectives in combination
ATC code: S02CA02

Mechanism of action
Flumetasone pivalate is a moderately potent glucocorticoid designed for local application. It exerts an anti-inflammatory, anti-allergic, vasoconstrictive, and anti-proliferative effect.

Experimental results show that the multiple effects of glucocorticoid drugs can be attributed to a complex molecular mechanism that includes binding to specific cytoplasmic receptors.

Pharmacodynamic effect
In inflammatory skin diseases of the external auditory meatus it affords prompt relief and eliminates symptoms such as pruritus while at the same time reducing swelling.

Appropriate studies in various animal species demonstrated the anti-allergic and anti-anaphylactic properties of LOCORTEN-VIOFORM® eardrops, solution.

Clinical efficacy and safety
Clioquinol, the antimicrobial component of LOCORTEN-VIOFORM® ear drops is a halogenated quinolinol derivative and is active against a broad spectrum of pathogenic micro-organisms, including fungi (e.g. Candida albicans and dermatophytes, Microsporum, Trichophyton) and gram-positive bacteria (e.g. staphylococci and streptococci).

Clioquinol has only a moderate inhibitory effect on gram-negative bacteria.

Clioquinol exerts a bacteriostatic, rather than a bactericidal action.

Microbiological testing demonstrated the antimicrobial and antimycotic activity of clioquinol, which is not influenced by the presence of the corticosteroid.

Pharmacokinetic properties

No pharmacokinetic data on LOCORTEN-VIOFORM® ear-drops are available.

Absorption
Trials (including treatment under occlusive dressings) with different formulations of LOCORTEN-VIOFORM® for topical application have shown that no demonstrable percutaneous absorption of flumetasone pivalate occurs, while clioquinol was absorbed to an extent of about 1.5% to 4%, as judged by the urinary excretion.

Elimination
Investigations of different pharmaceutical forms for topical use have shown that clioquinol is is absorbed and excreted in the urine mainly in glucuronide form and to a smaller extent as sulphate, whereas unchanged clioquinol is found in traces only.

Preclinical safety data

Teratogenicity
Some topical corticosteroids have been shown to be teratogenic in animals. The specific experimental teratogenic potential of glucocorticoid drugs in animals is known.

Mutagenicity
Flumetasone pivalate was not tested for mutagenic potential. Information on the mutagenicity of other corticosteroids suggests that this class of drug has minimal mutagenic potential. Clioquinol was shown to be non-mutagenic in studies using *Salmonella typhimurium*.

Carcinogenicity
Flumetasone pivalate has not been tested for carcinogenic potential. Another related topically-applied corticosteroid, fluticasone propionate, was negative for carcinogenicity following oral and topical administration. Clioquinol has not been tested for carcinogenicity.

**Pharmaceutical Particulars**

**List of excipients**
Polyethylene glycol 300 base

**Incompatibilities**
None known to date.

**Shelf life**
3 years

**Special precautions for storage**
Store below 25°C

**Nature and contents of container**
Yellow coloured polyethylene 10ml bottles containing 7.5ml solution.

**Instructions for use/handling**
Medicines should be kept out of the reach of children.

**Medicine classification**
Prescription Medicine

**Name and address**
AFT Pharmaceuticals Ltd
PO Box 33 203
Takapuna AUCKLAND
Telephone: 09 488 0232
Email: customer.service@aftpharm.com

**Date of preparation**
September 2016