CICLOSPORIN EYE PREPARATIONS FACT SHEET
TREATMENT OF OCULAR INFLAMMATORY CONDITIONS

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2. v1.1 – comments from CCG – approved at NCL JFC [26.03.15]
3. v1.2 – updated factsheet to include reference to the licensed product (Ikervis®, Santen Pharmaceuticals) [03.08.15]
4. V1.3 – Updated to include NICE TA369. Off label indications specified. [10.03.16]
5. V1.4 – “Ocular Rosacea” changed to “blepharokeratoconjunctivitis” (BKC) [16.08.16]

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Ciclosporin Eye Preparations
FACT SHEET - FOR THE TREATMENT OF OCULAR INFLAMMATORY CONDITIONS

Ciclosporin eye preparations should only be initiated in secondary care by an ophthalmologist specialising in corneal/external eye diseases.

Check List and Actions for GPs

- Practices must ensure that the information sheet / letter has been received from an ophthalmologist with indication for use, likely duration of treatment and evidence that the specialist has discussed with the patient that the product may be used for an off-label indication / dose (where relevant)
- Practices must ensure that the patient meets criteria for continuation of treatment (prescription initiated by an ophthalmologist specialising in corneal/external eye diseases and patient stabilised on treatment)
- Practices must ensure that repeat prescriptions will guarantee continuation of treatment until review

NICE

NICE state that ciclosporin (Ikervis) is recommended as a possible treatment for people with dry eye disease that has not improved despite treatment with artificial tears. [https://www.nice.org.uk/guidance/ta369](https://www.nice.org.uk/guidance/ta369)

Initiation

Ciclosporin eye preparation may be started in patients with the indicated diseases who have been found to be dependent on topical steroid to keep them free from sight-threatening ocular surface inflammation and/or free from severe and debilitating symptoms. No special tests or results are required before initiating treatment

Products

Ciclosporin eye preparations may be recommended as a treatment option after first-line agents (e.g. ocular lubricants / steroids) have failed or are not tolerated for:

- Atopic Keratoconjunctivitis (AKC) & Vernal Keratoconjunctivitis (VKC)
- Dry Eye Disease (DED)/ Keratoconjunctivitis Sicca (KCS)
- Blepharokeratoconjunctivitis (BKC) / Ocular Rosacea
- Thygeson’s keratitis & Chronic graft versus host disease

Ikervis

- Ciclosporin 1mg/ml eye drops [emulsion] – single use
- Licensed for the treatment of severe keratitis in adult patients with dry eye disease, which has not improved despite treatment with tear substitutes
- One ml of emulsion contains 1mg of ciclosporin and 0.05mg cetalkonium chloride
- Ikervis is supplied in 0.3ml single-dose containers presented in a sealed laminate aluminium pouch (one pouch contains five single-dose containers)
- Each single-dose container is sufficient to treat both eyes. Any unused emulsion should be discarded immediately
- NHS indicative price £72.00 30unit doses (April 2017)
Ciclosporin 0.2% eye ointment (unlicensed)
- 3.5g tube containing ciclosporin 2.0 mg/g
- Preservative free
- Drug Tariff price £76.40 (April 2017)

Choice of ciclosporin eye product
- The licensed ciclosporin eye product (currently Ikervis) should be used first line
- Some patients may not tolerate ciclosporin 1mg/ml eye drops (Ikervis) due to sensitivity to excipients. In such patients it may be necessary to prescribe ciclosporin 0.2% eye ointment
- Ciclosporin 0.05% eye drops (Restasis) should not be prescribed
- Ciclosporin 2% and 0.06% eye drops (unlicensed specials) are no longer manufactured and should not be prescribed

Administration
- Dosage: The recommended dose is one drop of IKERVIS once daily to be applied to the affected eye(s) at bedtime. Occasionally more frequent administration may be required at the discretion of consultant for recalcitrant disease. Duration of therapy: Subject to the patient diagnosis. To be communicated in GP letter. Ciclosporin 0.2% eye ointment may be used up to four times daily. Patients are to be reviewed at least every 6 months

Therapeutic Drug Monitoring
- Therapeutic drug monitoring is not required for topically administered ophthalmic ciclosporin products

Storage
- Ciclosporin eye products should be stored below 25°C. Do not freeze
- Ikervis: After opening the aluminium pouches the single-dose containers should be kept in the pouch in order to protect from light and avoid evaporation.
- Shelf life from manufacture: 3 years (Ikervis) / 2 years (ciclosporin 0.2% eye ointment)
- In use expiry (Ikervis): opened individual single-use containers with remaining emulsion should be discarded immediately after use

Adverse drug reactions (ADRs) reported
- Side effects from therapy are likely to be localised, transitory and mild to moderate in severity. The most common side effects reported were eye pain (19%), eye irritation (17.8%), lacrimation (6.2%), ocular hyperaemia (5.5%) and eye lid irritation (1.7%)

Associated risk
- Ikervis contains cetalkonium chloride which may cause eye irritation
- Contact lens wear should be avoided unless under specialist advice. If the eye drops are being instilled once daily at bedtime, the contact lenses should be removed before instillation of the eye drops and may be reinserted after waking-up the next morning

Procurement
- Ikervis ciclosporin 1mg/ml eye drops may be obtained by community pharmacies from usual commercial wholesalers
The annual cost of treatment with Ikervis is £864 based on frequency of once daily administration.
Ciclosporin eye ointment may be obtained from veterinary wholesalers.
## Appendix One: Quick Reference Guides

### Ciclosporin 1mg/ml eye drops [emulsion] (Ikervis®)  
Quick reference guide

<table>
<thead>
<tr>
<th><strong>Product name &amp; form:</strong></th>
<th>Ciclosporin 1mg/ml eye drops [emulsion] (Ikervis®)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Presentation:</strong></td>
<td>Single-dose, single-use containers</td>
</tr>
<tr>
<td><strong>Category:</strong></td>
<td>Licensed medicine</td>
</tr>
</tbody>
</table>
| **Indications of Use:**  | 1. Atopic Keratoconjunctivitis (AKC) & Vernal Keratoconjunctivitis (VKC) (Off label but approved for use in primary care by NCL JFC)  
2. Dry Eye Disease (DED)/ Keratoconjunctivitis Sicca (KCS)  
3. Blepharokeratoconjunctivitis (BKC) / Ocular Rosacea (Off label but approved for use in primary care by NCL JFC)  
4. Thygeson’s keratitis & Chronic graft versus host disease (Off label but approved for use in primary care by NCL JFC)  
5. Severe allergic eye disease (Off label but approved for use in primary care by NCL JFC) |
| **Rationale for Use:**   | Previous alternatives have failed to treat underlying condition |
| **Dose:**                | Normally once daily at night                      |
| **Duration of Treatment:** | Long-term (specific duration dependant on patient diagnosis) |
| **Cautions:**            | • Wash hands after use  
• Do not use in pregnancy  
• Do not use when there is fungal disease of the eye |
| **Side effects:**        | Mild irritation in the first few days of treatment may occur |
| **Monitoring:**          | No specific monitoring required as systemic absorption is minimal |
| **Storage:**             | Store below 25°C. Do not freeze. Discard unused content of each container immediately after use |
| **Other Information:**   | • Licensed for the treatment of severe keratitis in adult patients with dry eye disease, which has not improved despite treatment with tear substitutes  
• Contains cetalkonium chloride, medium chain triglycerides, glycerol, tyloxapol, poloxamer 188, sodium hydroxide (to adjust pH) and water for injection |
| **Ordering Information** | Directly from commercial wholesalers |
Ciclosporin 0.2% eye ointment
Quick reference guide

<table>
<thead>
<tr>
<th>Product name &amp; form:</th>
<th>Ciclosporin 0.2% eye ointment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Presentation:</td>
<td>3.5g eye ointment</td>
</tr>
<tr>
<td>Category:</td>
<td>POM-V</td>
</tr>
</tbody>
</table>
| Indications of Use (all unlicensed but approved for use in primary care by NCL JFC): | 1. Atopic Keratoconjunctivitis (AKC) & Vernal Keratoconjunctivitis (VKC)  
2. Dry Eye Disease (DED)/ Keratoconjunctivitis Sicca (KCS)  
3. Blepharokeratoconjunctivitis (BKC) / Ocular Rosacea  
4. Thygeson’s keratitis & Chronic graft versus host disease  
5. Severe allergic eye disease |
| Rationale for Use:            | Previous alternatives have failed to treat underlying condition |
| Dose:                         | Up to four times daily        |
| Duration of Treatment:        | Long-term (specific duration dependant on patient diagnosis) |
| Cautions:                     | • Wash hands after use  
• Do not use in pregnancy  
• Do not use when there is fungal disease of the eye |
| Side effects:                 | Mild irritation in the first few days of treatment may occur |
| Monitoring:                   | No specific monitoring required as systemic absorption is minimal |
| Storage:                      | Store below 25°C. Do not freeze. Once opened use within one month |
| Other Information:            | • Licensed as an animal medicine but has been manufactured to exactly the same standards as required for licensed human medicines  
• Useful when patients cannot tolerate eye drops  
• Contains paraffin base, maize oil and lanolin alcohol |
| Ordering Information:         | Directly from veterinary wholesaler |
Appendix Two: Background and Clinical Information

Background
Ciclosporin A is a fungal antimetabolite used as an anti-inflammatory drug due to its ability to inhibit interleukin 2 activation of lymphocytes.\textsuperscript{1} It is used systemically to prevent graft rejection in organ and tissue transplantation as well as in the treatment of eczema and psoriasis.\textsuperscript{2} From the 1980s reports have highlighted that topical ciclosporin can be used to treat a variety of ocular inflammatory conditions including dry eye disease, vernal & atopic keratoconjunctivitis and ocular rosacea (blepharokeratoconjunctivitis (BKC)).\textsuperscript{3,4,5} Topical ciclosporin (CsA) also has niche applications in post adenovirus keratitis, Thygeson's keratitis and peripheral corneal melts (particularly Mooren's ulcer). Conventional long term treatment with topical steroids to reduce ocular inflammation risks severe adverse effects including cataract formation and increased intraocular pressure.\textsuperscript{2}

Historically, no licensed preparations have been available for use in the eye. However, various unlicensed preparations have been available from importers, veterinary manufacturers or specials manufacturers. The first and, at present only, licensed ciclosporin eye preparation is Ikervis.

Many patients have encountered difficulty upon trying to obtain topical ophthalmic ciclosporin in primary care due to a reluctance of GP's to prescribe. Such decisions may be based on a lack of information regarding choice of product, indication for use, safety, treatment duration, licensing status, cost, availability and place in therapy. This protocol aims to provide information to accompany informed prescribing.

Indications & Summary of Efficacy Data

1. **Dry Eye Disease (DED) / Keratoconjunctivitis sicca (KCS)**
   - DED is a multifactorial disease of the tears and ocular surface that results in symptoms of discomfort, visual disturbance, and tear film instability with potential damage to the ocular surface. It is accompanied by increased osmolality of the tear film and inflammation of the ocular surface.\textsuperscript{6} Use of lubricating eye drops does not directly address the ocular surface inflammation. Anti-inflammatory therapies should be considered for patients with moderate to severe disease with symptoms or evidence of corneal disease refractory to treatment.\textsuperscript{7} The role of ciclosporin in the treatment of dry eye was reviewed in the Report of the Management and Therapy Subcommittee of the International Dry Eye Work Shop (2007). This was a consensus document by a group of highly-respected experts on ocular surface disease, including one from the UK. The report concluded that anti-inflammatory therapies should be considered for patients with moderate to severe disease with symptoms or evidence of corneal disease refractory to treatment. As part of the literature search performed by Moorfields Eye Hospital, 12 trials were identified of which ten showed a demonstrable effect on dry eye. Further details of these trials can be obtained on request from the Moorfields Eye Hospital Medicines Management Team.

2. **Atopic Keratoconjunctivitis (AKC) & Vernal Keratoconjunctivitis (VKC)**
   - AKC is a bilateral inflammatory ocular disease associated with atopic dermatitis. AKC is chronic in nature with frequent corneal complications which can cause permanent loss of vision. Aetiology involves a complex immunomodulatory dysfunction including type I and IV hypersensitivity reactions with Th1 cytokine profile.\textsuperscript{8} Corneal complications are seen in up to 70% of patients and approximately 30% require corneal transplantation.\textsuperscript{9}
VKC is a seasonal chronic ocular allergic disease which is usually self-limiting. However in 5-30% of cases permanent ocular changes may occur with associated visual impairment.\textsuperscript{10} The immunopathogenetic mechanism is complex and involves an IgE-mediated immediate hypersensitivity response as well as a Th2 type immune reaction.\textsuperscript{11}

In both VKC and AKC it is not uncommon for patients to be dependent on topical steroid treatment to control their ocular surface inflammation. These patients are at significant risk of steroid-related ocular side-effects and this risk may be mitigated by use of topical ciclosporin to reduce or eliminate steroid-dependence.

González-López et al (2012) conducted a Cochrane review on topical ciclosporin for atopic keratoconjunctivitis.\textsuperscript{12} The authors identified three RCTs with a total of 58 participants that were eligible for inclusion. One study reported on the use of 2% CsA in maize oil and two on the use of a commercial emulsion of 0.05% CsA. Of these three studies, one showed a beneficial effect of topical CsA in controlling signs and symptoms of AKC,\textsuperscript{13} one in controlling signs of AKC\textsuperscript{14} and one did not show evidence of an improvement\textsuperscript{15}. Only two studies analysed the effect of topical CsA in reducing topical steroid use; one showed a significant reduction in topical steroid use with CsA\textsuperscript{14}, but the other did not show evidence of this improvement.\textsuperscript{15} No serious adverse events were reported in the trials.\textsuperscript{13,14,15} The authors concluded that topical CsA may provide clinical and symptomatic relief in AKC and may help to reduce topical steroid use in patients with steroid-dependent or steroid-resistant AKC. The authors also remarked that no serious adverse events were identified in the trials.

A recent systematic review and meta-analysis of topical ciclosporin in allergic conjunctivitis reviewed results from 3 trials in AKC and 4 trials in VKC and concluded that the topical ciclosporin was more effective than placebo in alleviating the overall signs and symptoms of allergic conjunctivitis. However, the clinical and methodological heterogeneities in the studies mean that the overall efficacy should be interpreted with caution.\textsuperscript{16}

A literature search conducted at Moorfields Eye Hospital found 14 trials where topical ciclosporin was used for atopic or vernal keratoconjunctivitis, of which 13 showed a demonstrable effect. Further details of these trials can be obtained on request from the Moorfields Eye Hospital Medicines Management Team.

3. Ocular Rosacea / (blepharokeratoconjunctivitis (BKC))
Rosacea is an oculo-cutaneous inflammatory disorder which affects the facial sebaceous glands and the meibomian glands of the eyelid. Ocular rosacea is characterised by blepharo-keratoconjunctivitis, which can range from mild punctate epithelial erosions to significant corneal neovascularization and thinning. The skin shows erythema, telangiectasia and pustules over the cheeks, nose, and forehead with thickening over the nose (rhinophyma) in the later stages of this potentially sight threatening condition. The precise pathophysiology is unknown although a proposed theory is delayed type hypersensitivity like reaction. The traditional mainstay of treatment is similar to that of posterior blepharitis with topical antibiotics and steroids.\textsuperscript{7}

Van Zuuren et al (2011) conducted a Cochrane review on interventions for rosacea and concluded that ciclosporin 0.05% ophthalmic emulsion appears to be more effective than artificial tears for rosacea of the eyes.\textsuperscript{17} The authors identified ciclosporin as the only drug for which there was high grade evidence of effect in the disease, based on one RCT by Schechter et al (2009).\textsuperscript{18}
4. Niche applications in ocular surface inflammatory diseases

There are case series supporting the use of topical cyclosporine in a variety of chronic ocular surface inflammatory diseases such as Thygeson’s keratitis and chronic graft versus host disease.

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Disease</th>
<th>Trial characteristics</th>
<th>Number of patients (experimental / control)</th>
<th>Side effects</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reinhard T et al</td>
<td>1996</td>
<td>Thygeson’s keratitis</td>
<td>Case series, 2% CsA 3 times daily for a month with reducing regimen over 6 months</td>
<td>31 eyes of 17 patients</td>
<td>Burning and stinging</td>
<td>Complete resolution of opacities in 21/31 at 6 months. Partial resolution in a further 8/31.</td>
</tr>
<tr>
<td>Lelli GJ et al</td>
<td>2006</td>
<td>Graft versus host disease</td>
<td>Case series, 0.05% - 2% CsA 2-4 times daily</td>
<td>32 eyes of 16 patients</td>
<td>Irritation, burning</td>
<td>Corneal fluorescein staining improved in all patients. Dry eye symptoms improved in 10 patients (62.5)</td>
</tr>
</tbody>
</table>

Safety & monitoring

No significant systemic adverse effects have been reported in any of the trials found. Restasis has been used in the US since 2003. The most common adverse reaction reported on the Restasis prescribing information sheet is ocular burning (17%). Other reactions reported in 1% to 5% of patients included conjunctival hyperaemia, discharge, epiphora, eye pain, foreign body sensation, pruritus, stinging, and visual disturbance (most often blurring).21

Barber et al (2005) conducted a phase III, follow on, non-randomized safety evaluation of ciclosporin 0.1% ophthalmic emulsion, administered twice daily to dry eye disease patients. 412 adult patients, previously treated with ophthalmic ciclosporin, were treated for up to 3 years (mean duration of 19.8 months). The authors concluded ciclosporin was safe, well tolerated, and not associated with systemic side effects.22 The most common treatment-related adverse events were localised burning sensation (10.9%), stinging (3.9%) and conjunctival hyperaemia (3.4%).

The only other option open to patients with chronic inflammatory conditions are topical steroids which have a poor adverse effect profile, particularly in long term treatment. These adverse effects include cataract and raised intraocular pressure. When treated with topical steroids for 4–6 weeks 5% of the population demonstrated a rise in intraocular pressure greater than 16 mmHg and 30% demonstrated a rise of 6–15 mmHg.23,24 Patients using topical steroids require regular monitoring at 2-6 monthly follow up visits to screen for adverse effects whereas stable patients on CSA may be reviewed once per year.

Therapeutic drug monitoring is not required for topically administered ophthalmic ciclosporin products.

Supply

Initial supply will be made by the hospital pharmacy. Once stable, as judged by external / corneal ophthalmologist, the patient may obtain further supplies from primary care, subject to providing the GP with
relevant information containing indication, likely duration of treatment and evidence that a discussion with the patient has taken place regarding the unlicensed nature of the product.
Appendix Three: References