The 2016 Bowman Lecture
Conjunctival curses: scarring conjunctivitis 30 years on
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Conflict of Interest
I am involved with the development of ALDH inhibition for scarring under the Patent PCT/GB2015/051292 held by UCL Business plc.

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Abstract

This review is in two sections. The first section summarises thirty-five conditions, both common and infrequent, causing cicatrising conjunctivitis. Guidelines for making a diagnosis are given together with the use of diagnostic tests, including direct and indirect immunofluorescence, and their interpretation. The second section evaluates our knowledge of ocular mucous membrane pemphigoid, which is the commonest cause of cicatrizing conjunctivitis in most developed countries. The clinical characteristics, demographics and clinical signs of the disease are described. This is followed by a review and re-evaluation of the pathogenesis of conjunctival inflammation in mucous membrane pemphigoid (MMP), resulting in a revised hypothesis of the autoimmune mechanisms causing inflammation in ocular MMP. The relationship between inflammation and scarring in MMP conjunctiva is described. Recent research, describing the role of aldehyde dehydrogenase (ALDH) and retinoic acid (RA) in both the initiation and perpetuation of profibrotic activity in MMP conjunctival fibroblasts is summarised and the potential for antifibrotic therapy, using ALDH inhibition, is discussed. The importance of the management of the ocular surface in MMP is briefly summarised. This is followed with the rationale for the use of systemic immunomodulatory therapy, currently the standard of care for patients with active ocular MMP. The evidence for the use of these drugs is summarised and guidelines given for their use. Lastly the areas for research and innovation in the next decade are reviewed including the need for better diagnostics, markers of disease activity, and the potential for biological and topical therapies for both inflammation and scarring.

Keywords

conjunctival cicatrisation, conjunctival scarring, mucous membrane pemphigoid, autoimmunity, classification, diagnosis, pathogenesis, treatment
Introduction

It is a great privilege to have been asked to give this lecture and I am very grateful to the Scientific Committee of the College for inviting me to undertake it. I am going to discuss the results of 25 years of studies both collaborating with, and supported by, a large number of colleagues, including scientists and clinicians. I will use our studies in the context of the available evidence base to describe the causes and diagnosis of cicatrising conjunctivitis (CC) and follow this with a review of mucous membrane pemphigoid (MMP), which is the most common cause in developed countries. I hope that what I have to say about a group of diseases that is challenging for both patients and clinicians, will interest, entertain and challenge you as much as it has me.

All previous Bowman Lecturers have felt that their predecessors have been hard acts to follow and I am no different. However, all of us have worked in the shadow of Sir William Bowman. The portrait in Figure 1 has rarely been seen. It was painted by George Watts in about 1865 when Bowman was at the height of his powers. Watts was a friend of Bowman’s and the leading portraitist of the day. Like many others who have given this lecture I became fascinated by Bowman’s life and achievements. I wanted to know what it was that made him one of the foremost scientists of his day for his work on histology, bringing him Fellowship of the Royal Society at the age of 25, and then going on to become one of the founders of Ophthalmology as a scientific discipline in the UK. I have drawn on the biographies by his contemporary Power, and by James for the background, but also on other sources to make some connections that have not, to my knowledge, been made before. These have been summarised in Supplementary Appendix 1.

In the early 19th century the classification of conjunctivitis was almost unrecognizable from what we know today. However, by the end of the 19th century trachoma, trauma, ocular rosacea, adenovirus, allergic eye disease, lupus and pemphigus (at that time a term including all the pemphigoid diseases) had been recognized. A brief history of milestones in the recognition of the diseases causing CC is summarized in Supplementary Appendix 2 and Supplementary Table 1.

Current causes, epidemiology and classification of cicatrising conjunctivitis

Table 1 lists causes of CC from 2 studies. Thorne JE 2004 provides the largest institutional case series of CC cases attending an MMP clinic, either to have the cause of CC diagnosed, or to have ocular involvement by MMP excluded from those patients with an established diagnosis of extraocular MMP. But, causes in single clinics are skewed by case selection and referral bias. For this reason, the epidemiology of CC is best described
by the study from Radford C 2012 in the UK which lists the causes of cicatrizing conjunctivitis from the only national incidence study to have been reported. This study may have underestimated the incidence of CC by 15%, but the figures for MMP are in line with those reported for that disease from French and German estimates of 1.13 per million in France and 0.87 per million in Germany.

Table 2 lists 30 causes of CC with references and notes about the associated diseases and the rarity of the conditions. Although not complete, it includes those reported causes for which there is reasonable evidence of causation. These diseases have been divided into three categories in Table 2, each of which relate to current treatment protocols.

i Blinding scarring disorders include mucous membrane pemphigoid (MMP), and a rare subset of cases of drug induced scarring, and Stevens-Johnson syndrome/Toxic epidermal necrolysis (SJS/TEN), that behave like MMP and develop both inflammation and progressive scarring.

ii Ocular surface neoplasia (OSN) is an uncommon cause of inflammation and scarring which may mimic MMP although it is usually, but not always, unilateral unlike MMP which is usually, but not always, bilateral. The oncology protocols required for treatment will be delayed by misdiagnosis as CC.

iii Other conjunctival scarring diseases are those that have frequently been termed “pseudopemphigoid”. The reason for this distinction, and for the term pseudopemphigoid, has been summarized in the study from Thorne JE 2004 comparing MMP and with pseudopemphigoid where pseudopemphigoid is used as the term for non MMP causes of CC. “Other ocular surface diseases can cause cicatrizing conjunctivitis and symblepharon formation that mimic MMP. These diseases may be grouped under the term “pseudopemphigoid”. Although there are uncommon diseases associated with pseudopemphigoid that might require systemic immunosuppressive drug therapy (e.g., lichen planus or paraneoplastic pemphigus), most of the causes of pseudopemphigoid do not require systemic immunosuppressive drug therapy. Because long-term, systemic immune-suppressive drug therapy has potentially life-threatening side effects, differentiating ocular MMP from pseudopemphigoid is essential for the proper treatment of patients with cicatrizing conjunctivitis.”
However, the use of “pseudopemphigoid” as a catch-all term for any cause of CC that shares the clinical features of MMP, but which demands a different therapeutic approach, is too non-specific to be helpful for several reasons. It groups diseases with very different aetiologies, some of which may be associated with severe inflammation, unresponsive to topical therapy, that require immunosuppressive treatment. These include severe atopic keratoconjunctivitis (AKC), graft versus host disease (GVHD), sarcoid, ectrodactyly-ectodermal dysplasia-cleft syndrome (EEC syndrome) and ectodermal dysplasia (ED), even though the associated scarring rarely leads to pathology. Additionally, some of these conditions, with an established diagnosis, such as AKC and lupus, may also have concomitant MMP, which must be excluded. For these reasons the term pseudopemphigoid, as a specific term covering a disparate group of diseases, can be confusing and is best avoided.

Making the diagnosis

The diagnostic problem is shown by the similarity of the clinical appearances of the diseases illustrated in Figure 2. This has resulted in the difficulty we have in distinguishing many of the causes in Table 2 from ocular MMP and making a firm diagnosis, particularly for eye diseases without systemic associations or with dual pathology such as SJS/TEN, and for CC triggered by eye drops. Figure 3 summarises these dilemmas. Despite the similarities in the clinical characteristics some features of the clinical examination can help, as can a carefully taken history, and the appropriate investigations providing their limitations are understood.

Clinical examination, given the similarity of the clinical phenotype, is principally useful for establishing whether the disease is unilateral, and therefore potentially caused by ocular surface neoplasia, and to identify signs associated with a few of the other causes. Ocular MMP is rarely unilateral with only 7/115 (6%) of cases in one of our series. Other clinical findings that contribute are: severe aqueous tear deficiency (rare in early ocular MMP) caused by both Sjögren’s syndrome and severe non-Sjögren’s dry eye, for which a low Schirmer’s 1 test (without anaesthesia) is indicative although not diagnostic because of limited sensitivity. For ocular rosacea the scarring is associated with meibomitis and a papillary and follicular conjunctivitis with or without the following: pseudopterygium, peripheral corneal vascularisation and usually the dermatological features of acne rosacea.
History: The other disorders listed in Table 2 have a clear history of an associated ocular condition or systemic disease with the exceptions listed below. Direct questions about the eye diseases, risks associated with scarring, and systemic diseases that are associated with CC (listed in Table 2) will identify whether the patient has an ocular or systemic disease in association with CC. This information will not always be volunteered by patients or referring physicians. The diseases which develop CC, without a history of an associated systemic disease, or which present in the eye before disease develop elsewhere are:

- Ocular MMP (conjunctival disease without involvement of other sites) has comprised 32-48% of 132 cases in two UK studies including both cases with and without identified autoantibodies but meeting clinical criteria\(^4,39\).
- Paraneoplastic pemphigoid and pemphigus (PNP) may rarely present in the conjunctiva at the same time as a neoplasm, and before other signs of malignancy\(^27\).
- Pemphigus vulgaris (PV) may present in the conjunctiva before other signs: one case series of this presentation has been described\(^30\).
- Sarcoid may present in the conjunctiva and be associated with severe scarring\(^7\). Conjunctival biopsy, even in the absence of conjunctival signs, can be diagnostic in 30% of patients with systemic sarcoidosis\(^40\).

The rate of progression of scarring can usually be established by taking a careful history of the onset of symptoms of inflammation and scarring. The scarring may have been noticed by a healthcare professional or as a result of the onset of trichiasis. Estimating the rate of progression is important when deciding on the need for therapy, given the diagnostic delay for patients with ocular MMP (mean of 2.5 years and a range up to 10 years)\(^4\).

Patients with multiple causes of CC: as described in Table 2 patients with SJS/TEN and drug induced CC may rarely develop disease that is indistinguishable from ocular MMP. We also have one unreported case of MMP who later developed localised conjunctival OSSN. Other causes of CC including trachoma and atopic keratoconjunctivitis may also develop MMP and this can be expected to occur occasionally for any of the other causes. It follows that any patient with rapid progression of scarring and inflammation should have investigations to exclude MMP and OSSN.
**When to investigate?**

Unilateral cases (to exclude OSN), patients without a clear history of an associated ocular or systemic disease presenting in the eye (such as occurs in ocular MMP, PNP, PV and Sarcoid), and those with an ocular or systemic disease, but with inflammation and scarring not responding to topical therapy. The latter is most common in SJS/TEN, and in topical drug associated CC, for whom there is often reluctance to discontinue topical therapy for advanced glaucoma. In the latter group severe inflammation and scarring may be associated with most non preserved glaucoma medications, as well as due to preservatives\(^1\): the inflammation will often rapidly resolve on withdrawal of the drops although this may take 2-6 weeks to improve and longer to resolve\(^1\). Oral acetazolamide will often be tolerated by this group of patients, at least for 1-2 weeks to confirm the diagnosis of toxicity/allergy to topical medications. If oral acetazolamide is not tolerated, topical unpreserved iopidine, to which few of these patients have been exposed, can be used as an alternative. We have had to carry out glaucoma tube surgery on some patients having these reactions.

**Which investigations to use?**

Our protocol for the investigation of CC cases is included in Supplementary Appendix 3.

**Blood tests** are used in our department for the following:

a. Indirect immunofluorescence and individual autoantibody detection to identify specific autoantibodies associated with MMP and PV.

- **Indirect immunofluorescence**: Indirect immunofluorescence (IIF) microscopy on salt-split skin is used to identify the presence of serum autoantibodies in pemphigoid diseases. The structure of the basement membrane, its constituent proteins and the identification of these by immunofluorescence tests is illustrated in Figures 4 and 5\(^9\). However, IIF is non-specific for individual antibodies which cannot be identified from this test. The technique is clearly described by Hintner\(^42\).

- **Autoantibody detection**: autoantibodies to BP180, laminin 332, BP230, α6β4 integrin, laminin 311, type VII collagen, Laminin γ1 (p200 protein) are associated with MMP and those to the intraepithelial proteins desmoglein 1 and 3 are associated with pemphigus vulgaris\(^43\) which may occasionally present in the eye. ELISAs for antibodies to BP180, BP230, Collagen VII, Desmoglein 1 and Desmoglein 3 are
available from immunodermatology services and some laboratories offer immunoblotting to detect
the remaining autoantibodies.

Although often negative in MMP, positive indirect immunofluorescence, or autoantibody
detection, showing circulating anti-basement membrane zone antibodies provides additional evidence
for an underlying autoimmune pathology and may occasionally be positive when DIF is negative.

b. Autoantibody screening, and inflammatory markers to identify patients at risk of subclinical autoimmune
disease which is present in over 30% patients with MMP (compared to less than 10% in a control
population).

c. Baseline screening for patients who might need immunosuppressive therapy: full blood count, blood film,
glucose 6 phosphate dehydrogenase (G6PD) level (dapsone is contraindicated if this is elevated, which is
common in Mediterranean races), thiopurine methyltransferase (TPMT) levels relevant to metabolism of
azathioprine, urea and electrolytes, creatinine, liver function tests, glucose and infection markers including
quantiferon gold (for TB), hepatitis and HIV.

Biopsies are taken for routine histopathology and direct immunofluorescence studies. Bulbar conjunctival
biopsies are safe to take in patients with probable or proven MMP: of 344 cases having bulbar conjunctival
biopsy no adverse effects were reported. However, Foster reported that fornix conjunctival biopsies may
be associated with an exacerbation of scarring in 3 cases, following which he abandoned this site for diagnostic
biopsy. The technique for taking biopsies has recently been described.

a. Routine histopathology with fixation in 10% formol saline is used to identify atopic keratoconjunctivitis,
sarcoid and is essential to exclude OSN, particularly in unilateral CC.

b. Direct immunofluorescence (DIF) from perilesional tissue is required to confirm the diagnosis in the
immunobullous diseases. Figure 5 illustrates the differences between the DIF findings for the
intraepithelial “pemphigus” diseases and the subepithelial “pemphigoid” diseases, first defined by Lever in
1953. Lichen planus, an inflammatory disease affecting skin and sometimes the oral and ocular mucosa,
is also shown. DIF is used to differentiate these conditions and is described in Figure 5.

Interpretation of DIF results
Causes of cicatrising conjunctivitis which may be DIF positive from the conjunctiva and skin, or other mucosal sites, include:

- MMP (including mucosal dominated EB. Linear IgA disease and anti-laminin 332 pemphigoid), some patients with SJS/TEN, some with drug induced scarring, all of which show IgG or IgA and complement on DIF at the epithelial basement membrane (shown in Figures 4 and 5), should be treated as MMP cases.

  Note that for the diagnosis of ocular MMP a conjunctival biopsy is NOT required for the diagnosis if there has been a positive biopsy from any other site (skin, buccal, genital, nasopharyngeal mucosa), providing the ocular findings are typical of MMP. The DIF findings in BP are the same as for MMP so that the diseases must be differentiated on clinical grounds.

- Pemphigus vulgaris may rarely present in the conjunctiva as conjunctivitis in addition to conjunctival scarring, and conjunctival DIF has shown the characteristic epithelial intercellular fluorescence to IgG and complement shown in Figure 5, that is typically found in the skin at the dermoepidermal junction (basement membrane zone) in PV.

- Lichen planus has a characteristic “ragged” fibrin basement membrane deposition as shown in Figure 5.

Other diseases associated with CC, and which may have positive conjunctival DIF are:

- The ectodermal dysplasias with a pattern identical to that of MMP in the conjunctiva.

- Lupus shows granular deposition of IgG, IgA, IgM and complement at the dermal basement membrane in many cases and also in the conjunctiva in CC cases associated with lupus.

Conjunctival scarring associated disease which may be DIF positive from skin, but not from the conjunctiva:

- Conjunctival DIF has not been demonstrated in paraneoplastic pemphigus, which may also be associated with conjunctival scarring, and show a similar pattern to that of PV, in addition to a pemphigoid like pattern at the dermal epidermal junction.

- Dermatitis herpetiformis having a typical granular pattern at the basement membrane zone.

- Porphyria cutanea tarda also shows characteristic immunoglobulin deposition at the dermoepidermal junction but one study of associated conjunctival scarring showed repeatedly negative DIF results.
Negative immunofluorescence findings in ocular MMP

There is a strong recommendation, from an influential consensus document\textsuperscript{44}, that MMP can only be diagnosed when both clinical criteria and direct immunopathology criteria have been fulfilled. The clinical criteria are those distinguishing MMP from other diseases involving mucous membranes. Those that are listed in the document, and which cause CC, are lichen planus, SJS/TEN, PV, PNP, BP, and drug induced CC, to which the additional disorders summarized in Table 2 must be added. The direct immunopathology criteria that are recommended, as mandatory for diagnosis, are either direct immunofluorescence (DIF) microscopy, which is widely available, or immunohistochemistry which is not. Both positive IIF or specific autoantibody detection are widely accepted as alternative evidence when DIF findings are negative.

These criteria are usually positive in MMP that involves tissues other than the eye. However negative DIF results have long been recognized as presenting a diagnostic problem in patients with strong clinical evidence of ocular only MMP (in whom the conjunctiva is the only site of involvement). In three studies describing this group of 49 patients\textsuperscript{3,47,54} 25/49 (51%) were DIF+ of whom 13 (26%) required more than one biopsy to demonstrate this (Supplementary Table 2). Other problems with the use of DIF as a mandatory finding for the diagnosis of MMP are that the results can be initially positive, then subsequently negative when patients are in remission\textsuperscript{47}. Identical biopsy findings are found in bullous pemphigoid, which has to be distinguished by the clinical findings, and identical findings are also reported in patients some patients with SJS/TEN\textsuperscript{11}, drug induced progressive scarring\textsuperscript{55}, ectodermal dysplasia\textsuperscript{8} and ulcerative colitis\textsuperscript{54}.

Immunopathology, as described in the Consensus document, refers to the autoantibody detection tests (DIF, IIF or other means of autoantibody detection) that are recommended for diagnosis in these immunobullous disorders. However, the presence of autoantibodies does not reflect the dominance of the cell mediated autoreactive response, to mucosal basement membrane epitopes, which is probably important in many cases of ocular MMP (see sections b. and c. in the review of the pathogenesis of ocular MMP below). In effect a positive DIF result, as recommended for diagnosis, is a biomarker for an autoantibody driven autoimmune pathology and not the cellular autoimmune response. Unfortunately, DIF has not proved to be either sensitive in ocular MMP, nor specific as a marker for ocular MMP. Although a positive DIF result is useful, and can distinguish MMP from lichen planus, lupus erythematosus and PV and PNP, which have characteristic immune-pathological features of their own, a negative result does not exclude ocular MMP. Currently patients are being seen at Moorfields who could benefit from treatment for MMP, but who have been untreated.
because of uncertainty about the diagnosis due to their having negative autoantibody results. This has resulted in delayed therapy and deterioration of their scarring and inflammation. For these reasons we have proposed new criteria for the diagnosis of ocular MMP that are summarised below.

**Recommended diagnostic criteria for ocular MMP.**

We have proposed 3 sets of criteria for the diagnosis of ocular MMP:

- a. Patients with positive conjunctival DIF or positive DIF from another site meet currently agreed criteria.
- b. Patients with negative DIF from any site and positive indirect immunofluorescence can be diagnosed as having MMP.
- c. Patients with negative immunopathology can be diagnosed with ocMMP providing that they have a typical phenotype of progressive conjunctival scarring, and that other diseases that may cause this phenotype have been excluded. When ocular cases are reported, the detailed immunopathology findings should be recorded so that the diagnosis can be interpreted in light of future modifications to diagnostic criteria.

**When to refer?**

Patients with scarring & inflammation if there are no facilities for the specialist investigations or for delivering immunomodulatory therapies, for rarely described diseases, when the diagnosis is uncertain and when there is a poor response to therapy. In these circumstances a specialist centre with expertise in CC can provide the resources both to clarify the diagnosis and establish a treatment plan. Guidelines for referral are given in Figure 6.

**Review of mucous membrane pemphigoid with ocular involvement**

This is the commonest disease causing cicatrising conjunctivitis in the UK, accounting for 60% of CC cases, and is the prototypic autoimmune mucosal scarring disease. For the latter reasons MMP has been the focus of much of the research into scarring eye diseases in developed countries and ocular involvement by MMP has threefold more citations on PubMed than any other cause of CC apart from trachoma. MMP has been the focus of most of my work in CC and is the topic of the rest of this review.

**Terminology**
The terminology for the immunobullous diseases is confusing, given the changes in nomenclature over the past 250 years, and more recently. All these bullous diseases were categorized as pemphigus, until Lever’s seminal paper in which he differentiated the intraepithelial pemphigus diseases, from the subepithelial pemphigoid (meaning “resembling pemphigus”) diseases. This separation of these diseases was based on his histological studies demonstrating the tissue that was the focus of the inflammatory response. Since then the immunobullous diseases have been classified into these two groups, which are described in Figure 5.

The terminology for MMP has also changed. Initially Lever’s term for this was Benign Mucous Membrane Pemphigoid (BMMP) because, unlike pemphigus vulgaris, which had a very high mortality before the introduction of steroids, it was rarely fatal. In the 1980’s the term BMMP was replaced by the term cicatricial pemphigoid (CP) to reflect the associated scarring. CP and MMP have been used interchangeably at different times, and the studies describing the sites of involvement have prefaced these terms with the name of site, most often ocular CP (OCP) and ocular MMP (OMMP). Cicatricial pemphigoid is a term that is now used to describe a rare disease, having the same immunopathology as MMP, but which is limited to the head and neck and is also called localised CP or Brunsting Perry cicatricial pemphigoid. The term OCP is still in occasional use although, following the 2002 Consensus document the term MMP has been recommended for any patient, with any site of involvement (because multiple sites are often involved). However, this recommendation is impractical for specialist publications, describing the diagnosis and management of the different mucosal sites, and the descriptive terms ocular MMP (instead of OCP) and oral MMP are now being used for studies focused on diseases at these sites.

The other principal terminological difficulty is the altering nomenclature of the basement membrane immunoreactant proteins: examples are BP180, also known as Bullous Pemphigoid Antigen II and classified as Collagen type XVII, BP230 is also known as Bullous Pemphigoid Antigen I, laminin 332, formerly called laminin 5 or epiligrin, and p200 protein which is now designated as laminin γ1.

Clinical characteristics

a. Sites of involvement: Mucous membrane pemphigoid involves all the orificial mucosal sites (oral, ocular, nasopharyngeal, genital and anal) as well as, less often, tracheal and oesophageal. The skin may also be involved. Figure 7 shows examples of disease at many of the sites involved by MMP. All mucosal sites may become severely inflamed however scarring is rare in the oral mucosa whereas scarring is mandatory for
the diagnosis in the conjunctiva. Oral MMP, when restricted to the oral mucosa, has a relatively benign course in many patients, unlike ocular MMP. The conjunctiva is involved occurs in 70% of all MMP cases resulting in bilateral blindness in 20% of cases and severe sight loss in 30% of eyes. In a cross-sectional study (unpublished) between 2009-12 of 73 MMP patients with conjunctival involvement seen at Moorfields, 27% of patients had ocular only MMP, 26% ocular and oral, and a further 26% ocular, oral and nasopharyngeal, with the remainder having a mixture of other sites involved.

b. Demographics: In the UK national incidence survey which included 50 MMP patients meeting clinical criteria for the disease the median age was 71 years (range 20-90), there was a preponderance of males with a M:F ratio of 29:21 (1.38:1), with 32/50 (64%) having extraocular MMP (as opposed to ocular only MMP) of whom 8 (16%) had another autoimmune disease and 26/40 (65%) were taking immunosuppressive therapy at a 12 month follow up. These finding are similar to those in an institutional case series of 50 patients with a median age of 67 years (range 32-91), M:F ratio of 23:27 (0.85:1), extraocular MMP in 26/50 (52%) and 38/50 (76%) requiring immunosuppressive therapy after their initial assessment at the specialist centres. There is no good data on racial predisposition although MMP is probably less common in Indian Asians and the Chinese than in Caucasians. The disease is reported in Japan.

Clinical signs of disease progression

Early diagnosis and initiation of appropriate treatment are essential to prevent the sight-threatening complications of the approximately 75% of ocular MMP patients with rapidly progressive disease. Clinical signs are illustrated in Figure 8, ocular MMP typically presents with a red eye and persistent conjunctivitis that has not responded to topical therapy, or with cicatricial entropion and trichiasis, that may have failed surgical repair. About 30% of patients develop severe inflammation with limbitis leading to surface failure if uncontrolled of patients present with acute conjunctivitis and limbitis leading to rapidly progressive scarring and surface failure if uncontrolled. Similarly, persistent epithelial defect has a poor prognosis occurring in about 20% of patients. The remaining patients present with subacute or low grade chronic inflammation and slowly progressive scarring. The earliest clinical sign in patients with subacute disease is often medial canthal scarring, with loss of the plica and caruncle. Medial canthal scarring is usually an early sign of MMP and is not as frequent in conjunctival scarring due to other causes. Linear scarring in the sulcus subtarsalis (marginal
sulcus) of the upper tarsus is sometimes present early in the disease. Other signs, in order of progression, are
subepithelial reticular fibrosis, infiltration of the tarsal and bulbar conjunctiva, shortening of the fornices, symblepharon and cicatricial entropion, followed by ankyloblepharon and then, subsequent to scarring of the lacrimal ductules which usually occurs late in the disease, a totally dry “skin like” eye. Figure 9 describes the events leading to morbidity and blindness in ocular MMP.

Pathogenesis

a. Predisposing factors. As described in the section on the demographics of MMP above the disease is probably more common in Caucasians than in Indian and Chinese Asians, although it may occur in any racial group. Also, other autoimmune diseases are more common in MMP patients. Many patients have a genetic predisposition to MMP, expressing the HLA-DQB*0301 gene. For the majority of patients there are no identifiable precipitating factors. However, in subsets of patients, including cases of ocular MMP following SJS/TEN and topical glaucoma treatment, it is possible that damage to the conjunctival basement membrane precipitates the disease by exposing basement membrane epitopes triggering a pathological autoimmune response to neoantigens. The latter mechanism is an alternative to the development of loss of tolerance to basement membrane antigens that is described below and thought to be the underlying mechanisms in most cases.

b. Loss of tolerance, autoantibodies and autoreactive T cells and mechanisms of disease activation and remission at different sites in MMP. As in other autoimmune disorders disease probably develops as a result of loss of tolerance. In the case of the autoimmune subepithelial bullous dermatoses, of which MMP is one, loss of tolerance is to epithelial basement membrane proteins. This has been shown to result in circulating autoreactive T cells in 2 cases and in the generation of autoantibodies to a number of basement membrane proteins, most commonly BP 180, described in Figure 4. The pathogenic potential of antibodies has been demonstrated: anti-laminin 332 induces blistering in a mouse model and anti-α6β4 integrin induces separation of the dermoepidermal junction in organ culture of human skin. In serum, antibody levels have been correlated with disease activity in MMP. As a result of these findings it has been proposed that autoantibodies must be demonstrated for the diagnosis of MMP as for the other pemphigoid diseases.
On the other hand, not all patients with MMP have demonstrable circulating antibodies. In ocular MMP only 50% of patients have demonstrable autoantibodies (see section on negative immunofluorescence findings in ocular MMP, above). In addition, few MMP patients suffer from involvement of all potential sites of involvement: of 112 patients with MMP in a cross sectional study in London (unpublished) 2/112 patients had 5 involved sites, and 6/112 had 4 involved sites, from a potential maximum of seven sites (ocular, oral, skin, nasopharyngeal, laryngeal, genital, perianal). The mechanisms that protect individual sites from involvement in a systemic disease, caused by circulating antibodies to proteins common to the basement membranes of all the target tissues, may involve factors local both to those tissues and to the local cell mediated inflammatory response. The latter is a feature of these diseases in which autoreactive T cells are central both to the autoantibody response, and to the cellular autoimmune response. Combined cellular and antibody mediated responses are common in autoimmune diseases, all of which require autoreactive T cells. Although some autoimmune diseases are dominated by the pathogenic effects of one effector pathway, either autoantibodies or effector T cells, both pathways are commonly involved. Examples are Graves’ disease which is an autoantibody dominated disease, mediated by autoantibodies to thyroid stimulating hormone. At the other end of the spectrum, psoriasis is the result of an autoreactive T cell dominated response to skin associated antigens. However, most autoimmune diseases, exemplified by systemic lupus erythematosus and rheumatoid arthritis, result from the effects of both autoantibody and autoreactive T cell mediated inflammation in the target tissues. In bullous pemphigoid there is ample evidence for combined autoimmune cellular and antibody mediated effector pathways, as there is for mucous membrane pemphigoid. Autoimmune reactivity is normal in healthy individuals who may express both autoantibodies and autoreactive T cells without developing an effector response that results in disease. In the last decade naturally occurring autoantibodies have been shown to have homeostatic functions in the clearance of oxidatively damaged body waste, and in the modulation of immune cell functions. Autoantibodies in normal MMP controls: several studies have examined the prevalence of autoantibodies to BP180 and BP230, the basement membrane proteins commonly precipitating an autoreactive response in pemphigoid diseases and show that these are probably present in about 10-15% of healthy age matched individuals. However, the prevalence of these antibodies has varied in relation to the sensitivity and specificity of the assay used and the control group. The lowest prevalence has been than 1% in a large cohort of blood
donors of unknown ages, other publications have given values of 7.5%, 13%, 16% and 26% for ELISA and or immunoblots although higher values have been reported. Results for IIF have generally shown <5% positivity although this was 19% (n=32) in another study. Autoreactive T cells recognising the basement membrane protein BP 180, are also present in healthy individuals expressing the HLA-DQB*0301 gene (although not in those who did not). These autoreactive T cells are thought to be prevented from developing a pathological response by the activity of both regulatory T cells (Treg) and regulatory B cells (Breg). The role of these regulatory cells has been evaluated in only two studies in MMP: Treg were shown to be present in higher numbers in MMP lesional tissue compared to the skin of normal controls whereas a study of peripheral blood in patients, with a variety of pemphigus and pemphigoid diseases, showed reduced numbers of cells with Breg characteristics, compared to controls, in pemphigus only and not in pemphigoid.

In summary a substantial proportion of normal individuals have antibodies to both BP180 and BP230, some also having autoreactive T cells to BP180. There is evidence that Treg are present in MMP lesional tissue. We can hypothesise that differences in the balance between autoreactive T cells and Tregs in the different lesional MMP tissues account for the development and resolution of the disease in different target tissues in any one individual affected by MMP. If this occurs disease activation and remission in MMP is likely to be occurring at a local lesional level in the tissues, and independent of circulating autoantibody production and local antibody deposition. If this hypothesis is correct then the balance, between autoantibody and autoreactive cellular effector mechanisms, may differ both for different sites, and at different stages of disease chronicity, accounting for the lack of evidence of circulating antibodies in some cases of MMP, particularly in those with ocular MMP.

However, this is an area which demands further study, and the control mechanisms are likely to be more complex than this. Given that anti-CD20 B cell depletion therapy (such as the anti-CD20 monoclonal, rituximab) is effective in many cases of T cell mediated diseases such as type I diabetes and rheumatoid arthritis, as well as MMP, the role of B cells in regulating T cell responses has been examined and evidence suggests that, in some circumstances, B cell antigen presentation to autoreactive T cells may be necessary to develop an autoreactive T cell response. In addition, unstimulated B cells may also promote the development of Treg cells.
Summary of the pathogenesis of inflammation and scarring in ocular MMP.

This is described in Figure 10, for which the evidence is summarised both in Table 3 “Effectors and cytokines identified in MMP conjunctiva” and in the hypothesis synopsised above, that the inflammatory response is both the result of a variable balance between epithelial basement membrane autoreactive T cell mediated inflammation, in the lesional tissues, and also results from circulating autoantibodies to basement membrane proteins. Although the effects of inflammation and scarring are closely related it is easier to dissect out the mechanisms for each separately:

a. Inflammation: current descriptions ascribe the inflammatory response in MMP as arising from loss of tolerance to basement membrane proteins resulting in autoreactive T cells interacting with autoreactive B cells in the regional lymph nodes. This results in the generation of plasma cells, producing IgG or IgA circulating antibodies. Antibody may also be locally produced given that plasma cells are found in the lesional tissue, whereas the evidence for the presence of B cells in lesional tissue is mixed. Loss of tolerance is more common in individuals expressing HLA-DQB*0301 which may promote (restrict) the activation of T cells that are autoreactive to basement membrane proteins. Following this IgG and/or IgA antibody binds to the basement membrane in the conjunctival mucosa, resulting in complement fixation and the development of an acute inflammatory response leading to an influx of neutrophils, macrophages, dendritic cells and both cytotoxic (Tc) and helper (Th) T cells and plasma cells. MHC class II protein (required to present peptides to the immune system) are highly expressed in MMP conjunctiva. Mast cells and eosinophils, capable of producing pro-fibrotic cytokines are also found in acute disease.

Transforming growth factor beta (TGFβ) is overexpressed in acute disease compared to controls but not in chronic disease and, although the profibrotic factors, platelet derived growth factor (PDGF) and fibroblast growth factor (FGF) are present they are not overexpressed, suggesting that in chronic disease fibroblast activity remains functionally and morphologically abnormal after the withdrawal of the influence of growth factors. The B7-2 costimulatory molecule required for T cell proliferation in the presence of IL-2 is also overexpressed and can be expected to lead to increased T cell expansion. Of the cytokines, IL-2, IFNγ and TNFα (Th1 signature cytokines) are all present although not overexpressed compared to controls.

However, serum TNFα, an important pro-inflammatory mediator, has been shown to be elevated in MMP compared to controls and, in a more recent study, to be overexpressed in inflamed compared to treated...
MMP conjunctiva[^88]. Evidence for the activity of Th2 cells in ocular MMP comes from the finding of elevated levels of IL-13 in inflamed MMP conjunctiva[^89] and IL-5 in serum[^86], although the latter is also produced by mast cells.

b. **Scarring:** Profibrotic mediators have been investigated in several studies including TGFβ, PDGF, FGF and IL-13 as described above. Collagen type I and III are increased in ocular MMP stromal tissue. Cultured *in vitro* MMP conjunctival fibroblasts respond to TGFβ by the induction of heat shock protein 47 (HSP47), which is thought to influence procollagen synthesis and fibrosis resulting in collagen Type I production[^90].

Macrophage accumulation in the conjunctiva is probably an important event in the pathogenesis of MMP conjunctival scarring stimulated by macrophage derived cytokines and growth factors, including IL-4, PDGF, and TGFβ. Increased levels of macrophage inhibition factor (MIF)^[^91] and macrophage colony stimulating factor (m-CSF)^[^92], have been demonstrated in both *in vitro* cultured MMP fibroblasts and in whole conjunctiva and associated with increased numbers of macrophages. Connective tissue growth factor (CTGF), a downstream profibrotic mediator of TGF-β1, is overexpressed in both ocular MMP whole conjunctiva and in MMP cultured fibroblasts[^93]. These studies demonstrate that the effector cells, cytokines and growth factors necessary for fibrosis are present in the MMP conjunctiva. However, the mechanisms that relate this inflammatory milieu to the production of the extracellular matrix (ECM) by fibroblasts, that results in scarring, have not been identified[^94]. However, our recent studies outlined below have identified one control mechanism.

We have previously confirmed that OMMP fibroblasts maintain a profibrotic phenotype *in vitro* and have hypothesised that progressive fibrosis may be precipitated by the inflammation associated with ocular MMP, and then persist in eyes having clinical control of inflammation because of the continuing activity of persistently profibrotic fibroblasts[^97]. In recently published studies[^98] we have identified that the aldehyde dehydrogenase (ALDH)/retinoic acid (RA) metabolic pathway regulates this profibrotic activity in ocular MMP conjunctival fibroblasts *in vitro*. We have shown that ALDH is overexpressed in ocular MMP conjunctiva at the gene and protein level, compared to controls, and that ALDH inhibition with disulfiram abolished the profibrotic phenotype in MMP conjunctiva, resulting in the adoption of a normal control phenotype. Conversely *in vitro* fibroblasts from normal controls adopt a profibrotic phenotype when treated with RA, the metabolic product of ALDH. These findings provide evidence for ALDH/RA
autoregulation in ocular MMP fibroblasts as a mechanism underlying progressive conjunctival scarring seen in this disease and the potential for ALDH inhibition with disulfiram as a therapy for fibrosis. These findings were further confirmed in a mouse model of ovalbumin induced severe conjunctival inflammation that was developed for allergic eye disease studies in which we have shown that conjunctival scarring develops concurrent with inflammation. ALDH inhibition in this model, using topical disulfiram, was effective in preventing scarring in vivo, and also restored in vitro in mouse conjunctival fibroblasts to a normal phenotype, as in ocular MMP. Furthermore, another paper published with this study, has shown that in the same mouse model conjunctival scarring is initiated by the key role of dendritic cells, through paracrine production of ALDH/RA effecting conjunctival fibroblasts99. Given our hypothesis that the scarring in ocular MMP is the result of the inflammatory response in MMP, rather than due to the autoimmune pathogenesis per se, we believe this mouse model provides a surrogate for studying immune-mediated conjunctival scarring. Disulfiram is a drug already licensed for alcohol abuse control. These studies suggest that it could be repurposed for the topical treatment of conjunctival scarring in ocular MMP and provide justification for a randomized controlled trial of disulfiram therapy in this disease. Other mucosal scarring diseases are potential targets for further study using these techniques. Conjunctival diseases that may share these fibrotic mechanisms are Stevens-Johnson syndrome, atopic keratoconjunctivitis and trachoma.

Although ALDH inhibition is effective in these models we do not understand the molecular mechanisms underlying these profibrotic effects of ALDH/RA. It is possible that these profibrotic effects are mediated by similar mechanisms to those described in liver fibrosis 8,9, by the induction of the TGFβ1 gene, and/or activation of latent TGFβ1, for example by increased plasminogen activator levels. Active TGFβ drives critical changes in fibroblast metabolism, activation and ECM production in concert with pro-inflammatory cytokines and growth factors that affect fibroblast activity10. Alternatively, ALDH/RA might cause its effects through altered cellular energy metabolism via activation of TGFβ or as a consequence of the modulation of the transcription of metabolic genes11. The ALDH/RA metabolic pathway and these potential molecular mechanisms are described in Figure 10.

Treatment summary
a. Treatment outcomes for ocular MMP before 1980
These were poor until the 1980’s when immunomodulation techniques were introduced for their management for the first time. In the first series of cases reported by Morris in 1889 the outcomes were summarised: 28 cases (12F, 13M) onset from infancy to 76 years. “Disease began in the skin in 16, other mucosae in 4, and in the eye in 8. Twelve cases had conjunctival blistering, and others a pseudomembrane, entropion, progressive conjunctival shrinkage, cloudy cornea, thickened bulbar conjunctiva and xerophthalmia. Generally, vision is lost apart from perception of light although corneal perforation and destruction of the globe has been reported. One case had been described with spontaneous remission. Entropion surgery and application of lotions for the inflammation are palliative, there is no treatment for the progressive scarring. The pathology is obscure. It is a very rare disease”104. In Swanzy’s 1895 textbook he stated “Treatment is helpless in respect of arresting the progress of disease, or of restoring sight when lots in consequence of it. The most one can do is to relieve the distressing symptoms by emollients to the conjunctiva, and by the use of closely fitting goggles, to protect from wind, dust and sun. Internally arsenic is indicated105.” By 1951 Sorsby, in Systemic Ophthalmology stated of ocular pemphigus “Treatment is uniformly unsuccessful”106. The first textbook I owned when I started ophthalmology in 1978 was Parsons’ Diseases of the Eye 16th Edition in which it was stated that in the treatment of Benign Mucous Membrane Pemphigoid “Local treatment is unavailing as also, indeed, is general treatment”107. The use of dapsone for the management of pemphigus and pemphigoid was introduced in the 1970’s, but not for ocular MMP until 1982108. Systemic steroids for the control of acute disease in ocular MMP was described by Mondino in 1979109. The first use of immunosuppressive therapy, with azathioprine and cyclophosphamide, was described by Foster for 2 cases in 1980110, followed by a larger series using these drugs in 1982111. Subsequent publications have proliferated and are described below.

b. Current treatment

Successful management of disease demands the integration of the following:

(1) Control of surface disease

i. Blepharitis

ii. Dry eye

iii. Corneal punctate epitheliopathy

iv. Keratinisation
v. Trichiasis, entropion and lagophthalmos
vi. Persistent epithelial defects
vii. Corneal perforation
viii. Iatrogenic toxicity

(2) Control of immune mediated inflammation with systemic immunomodulation
(3) Control of fibrosis
(4) Prophylaxis of corneal ulceration and exposure
(5) Improving vision in patients with corneal blindness

The ocular surface disease management, and a synopsis of management with systemic
immunomodulatory therapy, and visual rehabilitation, has been discussed in detail in two reviews from
our group\cite{s1,s2} and a further recent review has summarized the use of immunomodulatory therapy in
detail\cite{s3}. In this publication I have restricted the description of treatments to a summary of the role of
immunomodulatory techniques and the evidence for their use in ocular MMP, with guidelines for their
use.

Use of systemic immunomodulation to control immune mediated inflammation

a. Lack of effect of topical immunosuppressive therapy in ocular MMP. Topical steroid treatment is ineffective
in controlling progressive ocular MMP, offering only variable symptomatic relief\cite{s4,s5}. Its adverse effects of
cataract and glaucoma generally outweigh the benefits. Subconjunctival steroids may be temporarily
effective, but relapses occur when the injections are stopped\cite{s6} and prolonged use also leads to cataract
and glaucoma. Anecdotally, topical steroid will relieve discomfort and inflammation in some patients with
mild/moderate disease activity but has not reduced the activity of severe inflammatory disease or the
progression of scarring. Topical ciclosporin has been used in only 4 reported cases of whom 2 had some
response\cite{s7}; we have little experience with this for ocular MMP and it is probable that the poor results
reported for systemic therapy with ciclosporin may have inhibited further investigation of this modality. As
a result, systemic immunomodulatory therapy is currently the standard of care for these MMP patients
\cite{s8,s9,s10}. 


b. Evidence for the effect of systemic immunosuppression on progression of disease. The primary goals of treatment of ocular MMP are to control inflammation and arrest fibrosis, in order to prevent progression of disease to more advanced stages and blindness. Most cicatrisation is occurs during active inflammation, but despite control of inflammation in 70% -78% of patients with systemic immunosuppression, progressive fibrosis was still observed in 61/115 (53%) and 23/54 (42%) of patients. Without treatment, conjunctival scarring in ocular MMP progresses in 13/20 (64%) of patients over 10 to 53 months. Progression is more frequent in the advanced stages of disease. Use of immunosuppressive therapy has been shown to slow progression of disease in one case series and control of inflammation has been shown to prevent progression in one RCT. With current immunosuppressive regimens, progression of cicatrisation has still been observed in 10 to 53% of ocular MMP patients and is more rapid in patients <60 years of age. A subset of MMP patients with ocular involvement have ongoing conjunctival fibrosis without overt clinical signs of inflammation. However, despite the absence of clinical signs of inflammation, there may still be significant cellular infiltrate on histological evaluation (“white inflammation”). Further systemic immunosuppression with potential systemic toxicity may not necessarily be helpful in these cases, for whom more specific local therapy targeting the cellular infiltrate or fibrogenic process would be ideal. There is evidence for both ongoing residual subclinical inflammation and transformed profibrotic fibroblasts as the putative drivers of scarring which progresses, despite apparent clinical control of inflammation with systemic immunosuppression. No current medical therapy is able to reverse the cicatrisation or ocular surface problems once they have developed.

c. Criteria for initiating immunosuppressive therapy. About 25% of OcMMP patients do not require immunosuppression as they have few symptoms, limited scarring, mild or no inflammation and slow progression, or are in remission. In end-stage “burned out” disease, immunosuppression is also unnecessary as eyes are usually comfortable, albeit blind, and treatment only slows scarring without significantly reversing it. In these patient groups, if conjunctival incision surgery is planned, such as that required for cataract extraction when the fornices are very short, or before fornix reconstruction, then immunosuppression should be started beforehand, to prevent an exacerbation of postoperative inflammation and scarring which may be severe, and will result in a poor surgical result and disease.
progression. For cataract surgery through a clear corneal incision, without conjunctival surgery\textsuperscript{123} or fornix incision for subtenon’s anaesthesia, and for lid surgery without a conjunctival incision, it is not necessary in my experience, to introduce preoperative immunosuppressive therapy.

d. Identifying inflammation due to underlying disease rather than to the secondary effects of a poor ocular surface. Identification of inflammation that is primarily due to the disease process in MMP, rather than to inflammation secondary to the effects of ocular surface disease or topical drug therapy, is essential before initiating immunomodulatory therapy. This is rarely a problem in those patients presenting with more severe disease. However, it can be difficult in those with less severe inflammation. It is necessary to differentiate inflammation due to the associated surface disease (dry eye, blepharoconjunctivitis, and lash abrasion), from immune mediated inflammation. Clinically this can be done by evaluating the degree of bulbar conjunctival inflammation under the upper lid, which is free of the worst effects of surface disease. However sectoral inflammation, although not common, does occur and may affect any quadrant of the bulbar conjunctiva as shown in Figure 11. Topical drop toxicity/allergy, typically resulting from glaucoma medications or preservatives may cause diffuse inflammation and be impossible to distinguish from that causes by MMP unless the topical therapy is discontinued: it takes 1-2 weeks for this to start to improve after withdrawing glaucoma medication, and any other topical drops, apart from non-preserved saline (see Section 3).

e. Evidence for the efficacy of different immunosuppressive regimens in controlling inflammation and guidelines for its delivery. Evidence for the effect of current immunosuppressive therapy in ocular MMP is summarised in Table 4 and comes from cohort studies\textsuperscript{124,125}, interventional and retrospective case series\textsuperscript{63,108,116,126-134} and two randomized trials\textsuperscript{46}. These have indicated a role for dapsone, sulfasalazine or sulphapyridine for mild to moderate inflammation, azathioprine, mycophenolate or methotrexate for moderate inflammation, or for disease not responding to sulphonamide therapy, and cyclophosphamide with a short course of prednisolone, for severe inflammation. Combinations of sulphas and myelosuppressive drugs (methotrexate, azathioprine, mycophenolate, and cyclophosphamide) with or without prednisolone can be effective. Notes on the use of these drugs, as “step up and step down therapy”, including combination therapy as favoured by UK specialists, are in Table 4 and Figure 12.
Supplementary Appendix 4 includes a synopsis of immunosuppression management guidelines for use in clinic.

f. Remission and when to stop therapy. Once complete control of inflammation has been achieved we continue immunomodulation, if well tolerated, for at least 12 months. Following this, the dose is reduced and can be stopped, if the patient wishes, providing they understand that it will need to be recommenced if disease activity recurs. Life-long follow-up is necessary, because disease recurs in up to 1/3 of patients. Our series of 115 patients showed remission without therapy of at least 6 months in 20 (17%) of patients with a relapse in 4/20 (20%). In our experiences long term remissions are uncommon using conventional drug therapies. However, Thorne et al. using high dose cyclophosphamide and prednisolone, achieved better remission rates, of at least 3 months without therapy, in 40/44 (91%) within 2 years of initiating therapy, followed by an 8/40 (20%) relapse rate. However, we think the risks of adverse effects outweigh the benefits as they require cumulative cyclophosphamide doses which probably exceed safe levels for the development of malignancy (see Table 4, rows 14 and 19).

The last thirty years and horizon scanning for the next decade

In the last 30 years there have been great advances in the understanding of the pathogenesis and therapy of three of the commonest causes of conjunctival scarring: trachoma, mucous membrane pemphigoid and atopic keratoconjunctivitis. These include the introduction of mass azithromycin delivery for trachoma control and the understanding that the scarring response in trachoma is only indirectly related to the precipitating infection resulting from the secondary profibrotic effects of infection. The unravelling of the pathogenesis of atopic keratoconjunctivitis and the randomised controlled trials, demonstrating the effectiveness of topical ciclosporin in its management. For ocular mucous membrane pemphigoid, the introduction of effective therapy with systemic myelosuppressives, currently the standard of care, had only just been introduced 30 years ago before which a majority of patients progressed to bilateral blindness. There was minimal understanding of the pathogenesis of MMP at that time and what we now know is feeding into the application of new therapies for inflammation. The latter include the use of monoclonal CD20 mediated B cell elimination, and the potential development of anti-scarring therapies. There are 4 areas in particular where advances can be expected to contribute substantially to improved outcomes for patients: improved diagnostic tests, the identification of
biomarkers of disease activity, the further development of biological drugs for therapy, and the development
of topical therapies for both inflammation and scarring.

a. **Better diagnostics**

The limitations of the current reliance on the detection of autoantibodies for disease identification,
particularly for ocular MMP, but also for MMP affecting other sites, have been outlined above. We
have some preliminary data using short peptides of BP180 that appear to be more sensitive than
currently available substrates for the identification of autoantibodies in ocular MMP. An alternative
approach has been used for the investigation of trachoma phenotypes using genetic biomarkers, that
are up and down regulated during the inflammation, scarring and resolution. A similar approach
in MMP, and other causes of CC, may be expected to provide an alternative route to autoantibody
detection, for both diagnosis and for the assessment of disease activity.

b. **Biomarkers of disease activity**

More objective biomarkers than clinical slit-lamp examination are needed to identify for both
inflammatory and scarring activity, to help identify the effects of therapy and guide their initiation and
withdrawal. Some progress has been made using neutrophils and their products for this. Expression of genes that are up and down regulated in inflammation and scarring may also be worth
investigating for this purpose.

c. **Biological drugs for MMP**

Conventional non biological immunomodulatory therapy with cyclophosphamide is effective anti-
inflammatory therapy from between 80% to 92% of cases in the larger series. However, the CD20
monoclonal Rituximab has been increasingly used as rescue therapy for patients with severe ocular
MMP who have failed cyclophosphamide therapy and with combined success rates of 80% in this
group. Although efforts have been made to carry out a randomised trial of this drug in ocular MMP
the cost of therapy has prohibited this, and manufacturers have been unwilling to support research
into an orphan indication. However, there are numerous CD20 monoclonal biosimilars in development
and it is probable that these drugs will become available for prospective case series, and randomised
controlled trials. The latter are needed to establish the effect of this drug in a number of areas. These include the effect in early ocular MMP, as opposed to unresponsive later stage disease. The length of remission, evidence for the induction of tolerance to autoantigens, and the identification of responders and non-responders needs to be investigated, as has been done in pemphigus\textsuperscript{162}. The optimum treatment regimen needs to be established, and the side effect profile compared to that of the other drugs used in this, relatively elderly, patient group having MMP.

d. Topical therapies for inflammation and scarring

There is an unmet need for effective topical therapies for inflammation and scarring to eliminate the morbidity associated with the side effects of systemic therapies. There are potential new topical therapies for both scarring and inflammation in CC. We have outlined the evidence for a promising new therapy, using ALDH inhibition, which has the potential to control scarring with topical therapy.

ALDH inhibition is also in development elsewhere as an anti-inflammatory therapy for allergic eye disease and anterior uveitis. Rapidly effective anti-scarring therapy for CC is critical in preventing morbidity, given the speed of scarring during severe inflammatory episodes which take 2-4 months to resolve with current anti-inflammatory therapies in ocular MMP, and which are also very rapid in diseases like Stevens-Johnson syndrome. The latter is another additional target disease for topical anti-scarring therapy which will be effective during acute phase inflammation, when most of the scarring occurs. Perfenidone is a new class of immunosuppressant with anti-inflammatory and antifibrotic effects. Its mechanisms of action are not fully established. In different model systems it has been shown to abrogate TGF-β1-stimulated collagen synthesis by inhibiting the upregulation of HSP47 and Col1 RNA, it blocks the proliferative effects of PDGF; reduces fibroblast proliferation and downregulates the proinflammatory cytokines, TNFα, IFNγ, IL-1B and IL-6\textsuperscript{163}. Topical application has been studied in human scleroderma\textsuperscript{164} and in glaucoma scarring models\textsuperscript{165}. Lastly, the topical use of anti-inflammatory monoclonal antibodies, such as the anti TNFα drugs which could be given subconjunctivally, should be evaluated.
Although a lot has been done in the last 30 years, the tools for understanding the pathogenesis of these diseases have expanded exponentially, and the potential for new therapies has seldom been greater, for patients cursed with these diseases.
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References


Legends for Figures

Figure 1

Portrait of Sir William Bowman by George Watts painted about 1865.
I am grateful to Rachel Clarkson, Sir William’s great-great-granddaughter, for providing this image.

Figure 2

Illustrations of some of the causes of cicatrising conjunctivitis

Figure 3

The diagnostic problem in cicatrising conjunctivitis
This Figure illustrates the classification of CC used in Table 2 with notes on the prevalence, which causes are often overlooked, which are usually known but which may co-exist with mucous membrane pemphigoid and causes associated with systemic diseases that may have their first manifestations in the eye.

Figure 4

The epithelial basement membrane structure and immunofluorescent tests
The cartoon describes the basement membrane (dermoepidermal junction) structure and its constituent proteins. The proteins that have been shown to be target antigens in MMP are listed. The position of the basement membrane (BM) is shown on a direct immunofluorescence (DIF) specimen which also shows positive immunofluorescence to IgG in the BM zone. An example of indirect immunofluorescence (IIF) on salt split skin is also shown demonstrating positive fluorescence to the floor (see Figure 5). The cartoon is reproduced with permission from Figure 1 Schmidt, E. & Zillikens, D. Pemphigoid diseases. Lancet 381, 320-332 (2013).

Figure 5

Autoimmune bullous dermatoses and lichen planus

Direct immunofluorescence (DIF) and indirect immunofluorescence
The illustrations in the top left panel show pemphigus vulgaris (PV) with DIF on perilesional skin showing intraepithelial antibody binding. In the top right panel, the skin lesions in bullous pemphigoid are shown for comparison with those in PV together with an example of a positive DIF in conjunctiva with antibody binding at the basement membrane. The pemphigus and pemphigoid diseases are listed separating those that may be associated with conjunctival scarring and those that are not. The panel on mucosal lichen planus shows the typical appearance of a “shaggy” fibrin staining band at the dermoepidermal junction. The bottom panel shows indirect immunofluorescence on human salt split skin from a patient with MMP.

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**Figure 5**

The spectrum of disease in mucous membrane pemphigoid (MMP)

- A. Gingival inflammation and ulceration
- B. Palatal inflammation and ulceration
- C. Supraglottic inflammation and scarring
- D. Oesophageal stenosis
- E. Skin ulceration and scarring
- F. Foreskin scarring
- G. Conjunctival inflammation and scarring
- H. Intraoperative photograph showing a normal subconjunctival space under incised severely scarred and shortened conjunctiva demonstrating that subconjunctival tissue is unaffected.

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**Figure 6**

Criteria for referral

Criteria for referral of patients with cicatrising conjunctivitis to a specialist ocular surface disease service. Adapted from Figure 5 in Williams, G.P et al. *Eye (Lond)* 25, 1207-1218 (2011).

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**Figure 6**

Ocular mucous membrane pemphigoid (MMP)

Clinical signs in ocular MMP

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**Figure 7**

Factors contributing to progression of disease in ocular mucous membrane pemphigoid (MMP)

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**Figure 8**
Summary of the pathogenesis of inflammation and scarring in ocular MMP

This illustrates the description in this section in the text, for which the evidence is summarised in Table 3. In brief a loss of tolerance to mucosal epithelial basement membrane proteins results in the development of pathogenic autoreactive T cells (aT) which help autoreactive B cells (aB) to proliferate in the regional lymph nodes, and differentiate into plasma cells. The latter produce circulating IgG and IgA autoantibodies to the mucosal basement membrane, which are detectable in the serum of some patients by indirect immunofluorescence (and other antibody specific assays). Plasma cells are also found in the conjunctival mucosal substantia propria where they may produce local antibody. In the mucosal epithelium direct immunofluorescence tests may show the presence of IgG and/or IgA fixed to the basement membrane where C3 (complement 3) is also identified in some patients. Activation of C3 at the basement membrane precipitates the complement cascade and acute inflammation at the basement membrane. This is the injury and inflammation phase of the disease causing an accumulation of inflammatory effector cells (neutrophils, dendritic cells, mast cells, eosinophils, macrophages and T cells) and the associated cytokines and growth factors IL-2, IL-5, IL-13 (interleukins), IFNγ, TNFα (tumour necrosis factor alpha), resulting in, often severe, inflammation and expansion of both Th1 and Th2 cells into a chronic inflammatory response. An alternative effector pathway may be more important, in the absence of autoantibody, in some subsets of patients whereby autoreactive T cells to basement membrane components home in on the mucosa. Here they can create an inflammatory response, in the absence of antibody, through their cytokines and growth factors including IFNγ, TNFα, IL-4, IL-5 and IL-13.

This inflammatory response results in fibrosis through the effects of profibrotic mediators released by macrophages, T cells, mast cells and eosinophils on fibroblasts, including PDGF (platelet derived growth factor), IL-13, TGFβ (transforming growth factor beta) and HSP47 (heat shock protein). Fibrosis also results from ALDH/RA (aldehyde dehydrogenase/retinoic acid) mediated paracrine effects of dendritic cells that activate a profibrotic phenotype in fibroblasts. During both active inflammation, and once inflammation has resolved, the activated MMP fibroblasts continue to scar as these remain profibrotic because of an ALDH/RA mediated autocrine effect. The latter probably results in RA dependent TGFβ activation and or induction, further driving fibrosis.

Figure 9
Sectoral conjunctival inflammation in ocular mucous membrane pemphigoid (MMP)

The same eye of a patient showing the superior bulbar conjunctiva free of inflammation but a localised area of inflammation in the inferior bulbar conjunctiva.

Figure 10

Immunosuppression for ocular mucous membrane pemphigoid (MMP)

This illustrates “step up” and “step down” therapy to control conjunctival inflammation as described in the section on Evidence for the efficacy of different immunosuppressive regimens in controlling inflammation and guidelines for its delivery and in Table 4 Row 34, and Saw VP 2008.

a. Drugs in different coloured boxes can be combined. The drugs in the dark red boxes (cyclophosphamide, mycophenolate, azathioprine and methotrexate) are all myelosuppressives: combining these results in unacceptable toxicity. Some authors have described the use of CD 20 inhibitors (such as Rituximab) with existing myelosuppressive therapy for unresponsive ocular MMP but this may be unsafe and we follow the recommendations of restricting adjunctive drugs to sulfas and stop the myelosuppressives immediately before the first CD20 monoclonal infusion. Prednisolone can generally be given for a short term effect with any of the other drugs listed but is usually needed in high doses, with its associated side effects.

b. In general, all of these drugs, except high dose steroids take 2-4 months to take effect. This period is probably shorter for the CD 20 monoclonals than the other drugs. A full effect will often take longer and for this reason, unless inflammation is deteriorating, I do not usually alter therapy for 12 weeks.

c. **For mild/moderate ocular MMP use step up therapy.** Start with a sulfa and, there is no effect, or toxicity, introduce mycophenolate. If the effect is limited but the drug is tolerated, add mycophenolate to the sulfa. Azathioprine (in grey typeface) is the second line drug when mycophenolate is not tolerated and methotrexate (in grey typeface) the third line drug in this situation because these are generally less well tolerated than mycophenolate and no more effective. If none of these drugs works switch to cyclophosphamide without using adjunctive steroid unless the patient is deteriorating. Failing these then a CD20 monoclonal is the next choice of drug and, if there is a lack of response to that 2 months after a second cycle of therapy, then start IV immunoglobulin therapy.
d. **For severe MMP** (severe conjunctival inflammation +/- limbitis +/- corneal ulceration) use **step down** therapy. Start oral Prednisolone 1 mg/kg WITH cyclophosphamide 1.0-1.5 mg/kg. Very severe cases will get a short term benefit from intravenous methylprednisolone 1 gram IV daily for 3 days before starting oral prednisolone with cyclophosphamide (unpublished data). If cyclophosphamide and prednisolone are not effective within 2-3 months, add a sulfa and then switch to a CD20 monoclonal. If this fails, as in the previous example, then start IVIG. Once control of inflammation has been established the patient can start stepdown therapy.

i. **For cyclophosphamide** follow the toxicity guidelines in Table 4 row 19, I usually maintain therapy for up to 12 months in the hope that a longer period of therapy might induce a remission. Two months before discontinuing cyclophosphamide, introduce a sulfa, if tolerated. After discontinuing cyclophosphamide patients with full control of inflammation can discontinue sulfas after another 2-4 months therapy. For patients with partial control the sulfa is maintained and mycophenolate (azathioprine or methotrexate) added. Drugs can be discontinued as required dependent on disease control and the development of side effects.

ii. After CD 20 monoclonsals patients may be in remission and any sulfa therapy can be discontinued after 2-4 months therapy. In the event of a relapse a decision has to be made whether to control this with step up therapy or a further cycle of a CD 20 monoclonal.

iii. After IVIG patients will usually still be in myelosuppressive agents which can be stepped down or stepped up as described above.
Figure 2

A MMP
B Trach
C Adeno
D Sebaceous
E OSSN
F EEC
G Eye drops
H AKC
I SJS
J Rosacea
K Lichen planus
L Sarcoid
Cicatrising conjunctivitis: all these causes may be clinically indistinguishable.

The Diagnostic problem

Blinding scarring diseases
Systemic immunomodulation needed

Surface Neoplasia
Usually unilateral scarring
Disease is usually known BUT MMP may need excluding

Other scarring diseases
Drugs
Eye drops: Glaucoma & others

Ocular diseases
Ocular rosacea
AKC
Trachoma
Other microbial Trauma

Systemic causes
Sarcoid
Other Pemphigoid
Pemphigus
Lichen planus
SJS/TEN
EEC
GVHD, Lupus Sjögrens

Mucous membrane pemphigoid

60% of all scarring conjunctivitis cases are MMP
Figure 4

Epithelial basement membrane structure and immunofluorescence tests

Epithelial basement membrane (BM)

Direct immunofluorescence (DIF)

Indirect immunofluorescence (IIF) on salt split skin: splits in the lamina lucida of the BM

Further examples of DIF and IIF are shown in Figure 5
Pemphigus is intraepithelial

Pemphigoid is subepithelial

Pemphigus diseases with ocular scarring
- Pemphigus vulgaris
- Paraneoplastic pemphigus

Pemphigus diseases with no ocular scarring
- Pemphigus foliaceus
- Pemphigus gestationis
- IgA pemphigus

Mucous membrane pemphigoid now also includes mucosal dominated linear IgA, epidermolysis bullosa acquisita, anti-laminin 332 pemphigoid (previously laminin 5 and epiligrin)
- Bullous pemphigoid

Pemphigoid diseases with ocular scarring
- Anti-p200/anti-laminin γ1 pemphigoid
- Lichen planus pemphigoides
- Pemphigus gestationis

Indirect immunofluorescence on human salt split skin
DIF on perilesional skin
DIF on conjunctiva
DIF on LP
Roof
Floor

Detects antibodies to BP180, BP230, and α6β4 integrin (on the “roof” of the split) and laminin 332, laminin γ1 and type VII collagen (on the “floor” of the split)
Cicatrising conjunctivitis: guidelines for referral to a specialist Ocular Surface Disease service

Do you have access to a specialist immunofluorescence diagnostic laboratory?  
No

Yes

Carry out:
(i) Supportive diagnostic conjunctival ± oral mucosal biopsies for Direct IF and serum for Indirect IF
(ii) Routine histology
• To exclude neoplasia in unilateral cases
• To assist in the diagnosis of atopic keratoconjunctivitis

Do you have expertise/access to delivering immunosuppression, patient counselling and an oculoplastics subspecialty service?  
No

Yes

Commence/continue appropriate management strategies and monitor patient

Has the conjunctival scarring worsened or has the eye become inflamed?  
No

Yes

Refer to Specialist Tertiary Hospital

Shared care
Spectrum of disease in mucous membrane pemphigoid
Loss of plica
Subepithelial fibrosis
Fornix shortening
Symblepharon
Entropion & trichiasis
Subepithelial fibrosis
Surface failure, keratin
Ulceration
Limbitis
PED
Ocular mucous membrane pemphigoid
PROGRESSION in ocular MMP

CONJUNCTIVAL SCARRING
- Entropion
- Trichiasis
- Lagophthalmos
- Dry eye

SURFACE DISEASE
- Punctate keratopathy
- Persistent defect & infection
- Corneal vascularisation
- Corneal scarring
- Keratinisation

IMMUNE MEDIATED INFLAMMATION
- Surface failure (stem cell failure) in severe cases
- Conjunctival overgrowth

SURFACE PAIN

CORNEAL BLINDNESS
**Injury phase & INFLAMMATION**

- **Loss of tolerance:** Autoreactive T cells recognise BMZ antigens
- **Antibody production in lymph nodes:** IgG or IgA
- **Complement** activated

**Acute inflammation**

- Mast cells
- Eosinophils
- Neutrophils
- Macrophage

**Chronic inflammation**

**SCARRING**

- Extracellular matrix production
- Fibrogenic factors
- T cell expansion
- T cell

**Direct immuno-fluorescence of MMP conjunctiva**

**Fibroblast**

- ALDH
- ALDH/RA effects
- RA dependent TGFβ activation/induction
Figure 11

Sectoral conjunctival inflammation in ocular MMP

Superior and inferior bulbar conjunctiva showing persistent sectoral inflammation in partially controlled disease
Immunosuppression for ocular MMP

Drugs in different colour groups can be combined

Wait for 12 weeks before switching

Oral prednisolone 1mg/kg (taper over 12 weeks) for rapid effect

CD 20 monoclonals

IV immunoglobulin

Cyclophosphamide

Mycophenolate

Azathioprine

Methotrexate

Mycophenolate

Azathioprine

Methotrexate

Dapsone or Sulphapyridine

For mild MMP

Dapsone or Sulphapyridine

Figure 12
### Table 1
Institutional and national surveys of cicatrising conjunctivitis

<table>
<thead>
<tr>
<th><strong>Institutional case series of cicatrising conjunctivitis (CC)</strong> Thorne JE 2004&lt;sup&gt;3&lt;/sup&gt;</th>
<th><strong>UK national survey of new cases of CC</strong> Radford C&lt;sup&gt;4&lt;/sup&gt; n=82</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MMP</strong></td>
<td><strong>Number (%)</strong></td>
</tr>
<tr>
<td></td>
<td>74 cases of MMP and 145 others</td>
</tr>
<tr>
<td><strong>Incidence per million</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Other causes of CC</strong></td>
<td><strong>Number (%)</strong></td>
</tr>
<tr>
<td>Linear IgA bullous dermatosis&lt;sup&gt;b&lt;/sup&gt; None (N)</td>
<td>1</td>
</tr>
<tr>
<td>Topical glaucoma drug induced (28.3%)</td>
<td>3</td>
</tr>
<tr>
<td>Rosacea blepharoconjunctivitis (20%)</td>
<td>3</td>
</tr>
<tr>
<td>Atopic keratoconjunctivitis (8.3%)</td>
<td>3</td>
</tr>
<tr>
<td>Sjögren’s syndrome or KCS&lt;sup&gt;c&lt;/sup&gt; (7.6%)</td>
<td>1</td>
</tr>
<tr>
<td>Stevens Johnson syndrome (6.3%)</td>
<td>16</td>
</tr>
<tr>
<td>Graft versus host disease (1.4%)</td>
<td>2</td>
</tr>
<tr>
<td>Lichen planus N</td>
<td>2</td>
</tr>
<tr>
<td>Ocular surface neoplasia N</td>
<td>1</td>
</tr>
<tr>
<td>Non-trachomatous infection (5.5%)</td>
<td>N</td>
</tr>
<tr>
<td>Trachoma (2.8%)</td>
<td>N</td>
</tr>
<tr>
<td>Sarcoidosis (3.5%)</td>
<td>N</td>
</tr>
<tr>
<td>Pemphigus vulgaris (3.4%)</td>
<td>N</td>
</tr>
<tr>
<td>Paraneoplastic pemphigus (2.8%)</td>
<td>N</td>
</tr>
<tr>
<td>Trauma or lid surgery (2.8%)</td>
<td>N</td>
</tr>
<tr>
<td>Ectodermal dysplasia (2.1%)</td>
<td>N</td>
</tr>
<tr>
<td>Chronic cutaneous lupus (1.4%)</td>
<td>N</td>
</tr>
<tr>
<td>Aplasia cutis congenital (0.7%)</td>
<td>N</td>
</tr>
<tr>
<td>Ectopic geographic tongue (0.7%)</td>
<td>N</td>
</tr>
</tbody>
</table>

<sup>a</sup>Percentages of causes for the 145 cases of cicatrising conjunctivitis not caused by MMP. Percentages total above 100% because some patients had more than one diagnosis.

<sup>b</sup>Now classified as MMP

<sup>c</sup>Keratoconjunctivitis sicca
Table 2
Current classification of cicatrising conjunctivitis.
Causes are underlined. Some causes appear twice because they result in either progressive or non-progressive scarring disease. The pemphigus and pemphigoid diseases NOT currently known to cause conjunctival scarring are also listed.

<table>
<thead>
<tr>
<th>CLASSIFICATION</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BLINDING SCARRING DISORDERS</strong></td>
<td>Progressive inflammatory and scarring diseases for which systemic immunomodulation is often needed for control of scarring &amp; inflammation</td>
</tr>
<tr>
<td>Mucous membrane pemphigoid (MMP)</td>
<td>Currently 60% of all CC cases in the UK are caused by MMP which now includes the conditions mucosal dominated epidermolysis bullosa acquisita, linear IgA disease and anti-laminin 332 (formerly anti-epiligrin or anti-laminin 5) pemphigoid</td>
</tr>
<tr>
<td>Drug-induced progressive conjunctival cicatrisation</td>
<td>A rare complication of topical medication, usually for glaucoma. A small subset of patients with these diseases develop autoantibody-positive or negative progressive conjunctival indistinguishable from MMP</td>
</tr>
<tr>
<td>Stevens-Johnson Syndrome (SJS) &amp; Toxic Epidermal Necrolysis (TEN) with progressive scarring</td>
<td>Only a small subset of patients with these diseases develop autoantibody-positive or negative progressive conjunctival scarring similar to that in MMP which may continue from the acute episode or develop acutely later</td>
</tr>
<tr>
<td><strong>OCULAR SURFACE NEOPLASIA</strong></td>
<td></td>
</tr>
<tr>
<td>Ocular surface squamous carcinoma (OSSN) and sebaceous cell carcinoma</td>
<td>Rare causes of conjunctival scarring and inflammation indistinguishable clinically from MMP, except that cases are usually unilateral</td>
</tr>
<tr>
<td><strong>OTHER SCARRING CONJUNCTIVAL DISEASES</strong></td>
<td>Diseases that may be clinically identical at presentation with fornix shortening &amp; symblepharon BUT in which scarring rarely results in blindness, although inflammation may be severe. MMP may occur in these diseases and should be excluded</td>
</tr>
<tr>
<td>Ocular diseases with no systemic involvement</td>
<td></td>
</tr>
<tr>
<td>Drug induced scarring</td>
<td>Usually due to topical preservatives or unpreserved glaucoma medication and, rarely, other drugs. Inflammation resolves after withdrawing the drops and scarring stabilizes</td>
</tr>
<tr>
<td>Atopic keratoconjunctivitis</td>
<td>Usually a tarsal papillary reaction in addition to scarring although this may be minimal in severe longstanding cases</td>
</tr>
<tr>
<td>Trachoma</td>
<td>Tarsal scarring typical but symblepharon occurs</td>
</tr>
<tr>
<td>Adenoviral conjunctivitis</td>
<td>Scarring with entropion and symblepharon are occasionally present. These patients must have a history of an acute severe conjunctivitis to confirm the diagnosis</td>
</tr>
<tr>
<td>Conjunctival trauma: chemical, thermal, surgical and radiation injury</td>
<td>Diagnosis is clear from the history</td>
</tr>
<tr>
<td>Oculodermal diseases</td>
<td></td>
</tr>
</tbody>
</table>
Stevens-Johnson syndrome and Toxic Epidermal Necrolysis

Diagnosis clear from history. A small subset of patients develop progressive scarring disease see the section on Blinding Scarring Disorders above.

| Pemphigoid diseases (sub-epithelial immunobullous diseases) | Pemphigoid diseases with conjunctival scarring unreported or very rare: one case of anti-p200 with anti-laminin 332 pemphigoid, lichen planus pemphigoides may involve oral but not ocular mucosa, pemphigoid gestationis rarely involves mucosa and conjunctival involvement is unreported to my knowledge.

| Bullous pemphigoid | Conjunctival scarring infrequent.

| Mucous membrane pemphigoid | See Blinding Scarring Disorders above. In about 20% of cases the disease progresses very slowly or is in remission and systemic immunosuppression is not required.

| Paraneoplastic pemphigoid | Severe scarring may occur.

| Pemphigus diseases (intra-epithelial immunobullous diseases) | Pemphigus diseases in which conjunctival disease is unreported: pemphigus gestationis, pemphigus foliaceous (may involve lid skin but not the conjunctiva) and IgA pemphigus.

| Pemphigus vulgaris | Conjunctivitis occurs but scarring is rare. However, several cases have presented in the eye accompanied by severe scarring and corneal perforation.

| Paraneoplastic pemphigus | Conjunctivitis occurs. Severe scarring may occur but is uncommon.

| Dermal diseases with variable ocular involvement | 

| Ocular rosacea | Some scarring is common.

| Dermatitis herpetiformis | Rarely causes scarring.

| Lichen planus | Rarely causes scarring.

| Systemic lupus and discoid lupus | Rarely causes scarring.

| Multisystem disorders | 

| Graft versus Host Disease | The diagnosis has usually been made by the oncologist. Conjunctival scarring and inflammation is common.

| Sarcoid | Rare but may present in the eye as severe scarring conjunctivitis.

| Sjögren’s syndrome and keratoconjunctivitis sicca | Scarring is rare.

| Miscellaneous congenital and acquired disease | 

| Ectodermal dysplasia (ED) diseases (includes ectrodactyly-ectodermal dysplasia-cleft syndrome (EEC) cases as well as ED) | Some patients develop circulating basement membrane antibodies or mucosal direct immunofluorescence: inflammation may be severe with scarring.

| Inflammatory bowel disease | Scarring is reported but very rare.

| Porphyria cutanea tarda | Scarring is reported but very rare.
Table 3
Effectors and cytokines identified in MMP conjunctiva

<table>
<thead>
<tr>
<th>GLOSSARY (alphabetical in the order of the abbreviations in the Table with additional notes on origin and function)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B7-2: a costimulatory T cell ligand expressed on dendritic cells and macrophages.</td>
</tr>
<tr>
<td>CD40, CD80, CD154: costimulatory molecules on T cells.</td>
</tr>
<tr>
<td>FGF: a family of fibroblast growth factors, including basic FGF (bFGF) produced by monocytes and important in wound repair.</td>
</tr>
<tr>
<td>CTGF: connective tissue growth factor produced by fibroblasts and is a downstream mediator of TGFβ1 induced collagen synthesis, promotes fibroblasts proliferation &amp; enhances matrix production.</td>
</tr>
<tr>
<td>HSP family: heat shock proteins produced by fibroblasts and involved in the maturation of collagen</td>
</tr>
<tr>
<td>ICAM: intercellular adhesion molecule expressed on macrophages and lymphocytes facilitating migration into inflamed tissues</td>
</tr>
<tr>
<td>IFNγ: interferon gamma, produced by activated T&lt;sub&gt;h&lt;/sub&gt;1 lymphocytes, epithelia and fibroblasts with multiple roles in inflammation.</td>
</tr>
<tr>
<td>IL-1: interleukin 1 produced by macrophages, B cells, lymphocytes, fibroblasts, and endothelium is a multifunctional cytokine critical in initiating and maintaining inflammation and immune responses.</td>
</tr>
<tr>
<td>IL-2: interleukin 2, produced by T cells, and responsible for T and B cells proliferation and activation.</td>
</tr>
<tr>
<td>IL-4: interleukin 4, produced by mast cells, Th2 cells and fibroblasts affecting B cell, T cell and fibroblast functions with multiple profibrotic and inflammatory effects. Induces m-CSF, type I collagen &amp; collagen-binding HSP47 production by conjunctival fibroblasts.</td>
</tr>
<tr>
<td>IL-5: interleukin 5, produced by mast cells, Th2 cells, eosinophils and monocytes and affects eosinophil chemotaxis, differentiation, activation and prolongs survival.</td>
</tr>
<tr>
<td>IL-6: interleukin 6, produced by macrophages, and stimulates B cells to differentiate into plasma cells</td>
</tr>
<tr>
<td>IL-13: interleukin 13, produced by Th2 cells, mast cells and basophils and is a potent stimulator of eosinophil, lymphocyte and macrophage rich inflammation and tissue fibrosis</td>
</tr>
<tr>
<td>m-CSF: macrophage colony stimulating factor produced by monocytes, fibroblasts and endothelial cells having effects on the proliferation, differentiation and activation of monocytes macrophages</td>
</tr>
<tr>
<td>MIF: macrophage inhibition factor produced by T cells and macrophages inhibiting macrophage migration and stimulating macrophage activation.</td>
</tr>
<tr>
<td>MPO: myeloperoxidase. This enzyme is produced and stored by neutrophils</td>
</tr>
<tr>
<td><strong>MMP</strong> (in italics to distinguish the abbreviation from that for mucous membrane pemphigoid) : matrix metalloproteinase family of enzymes including <strong>MMP-8</strong> and <strong>MMP-9</strong> that degrade extracellular matrix</td>
</tr>
</tbody>
</table>
- PDGF: platelet derived growth factor produced by macrophages and fibroblasts amongst others.
- Tc (Cytotoxic T cells) expressing CD8
- TGFβ family: transforming growth factor beta, produced by many cell types (B & T cells, monocytes, macrophages, fibroblasts). These are profibrotic mediators with immunosuppressive effects.
- Th1: T helper subset 1 expressing CD4 secreting their signature cytokine interferon gamma (IFNγ) and also producing TNFα.
- Th2: T helper subset 2 expressing CD4 and secreting the signature cytokine IL-4 and also producing IL-5 and IL-13
- TNFα: tumour necrosis factor alpha, produced by macrophages, mast cells and lymphocytes. It has multiple inflammatory functions

<table>
<thead>
<tr>
<th>Effector cells &amp; proteins</th>
<th>Findings in MMP</th>
<th>Author</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circulating autoreactive T cells to BP180</td>
<td>2/10 patients with MMP demonstrated IFNγ production to BP180 NC16A in peripheral blood mononuclear cells suggesting that this segment of BP 180 is a T cell target antigen in MMP</td>
<td>Black AP 200468</td>
</tr>
<tr>
<td>Neutrophils, macrophages, dendritic cells, Th (CD4), Tc (CD8), plasma cells. (TGFβ FGF and PDGF)</td>
<td>Conjunctival biopsies from 20 patients with acute (n4) subacute (n8) and chronic (n8) MMP investigated showed an excess of neutrophils, macrophages and dendritic cells in acute disease. Increased T cells present in all subsets, with Tc:Th ratio higher except in acute disease when these were equal. Plasma cells increased, but not B cells or NK cells. PDGF, FGF and TFGβ present in both diseased and controls. TFGβ was overexpressed in acute disease. MHC class II highly expressed in diseased tissue (potential to present antigen to Th cells). The activated T lymphocytes were identified with anti-IL-2 showing these were Th1 T cells.</td>
<td>Bernauer W 199385</td>
</tr>
<tr>
<td>Neutrophils in the epithelium</td>
<td>Neutrophils are elevated in the MMP conjunctival epithelium shown by impression cytology and increased levels are associated with progressive scarring in both inflamed and clinically uninfamed eyes and may be a useful biomarker of progressive fibrosis and response to therapy</td>
<td>Williams G 2016100</td>
</tr>
<tr>
<td>Neutrophil derived enzymes (MPO, MMP-8 and MMP-9)</td>
<td>MPO is neutrophil derived, and neutrophils are also a source of MMP’s which are elevated in the tears of MMP patients potentially accounting for their predisposition to corneal melting and also potentially of use as biomarkers of disease activity</td>
<td>Arafat S 2014101</td>
</tr>
<tr>
<td>Dendritic cells and</td>
<td>MMP epithelium and control did not differ. Substantia propria: predominantly</td>
<td>Tesavibul N</td>
</tr>
<tr>
<td><strong>T cell co-stimulatory molecules</strong></td>
<td>increased increased macrophages and T cells but also Langerhans &amp; B cells. B7-2 costimulatory ligand (usually expressed on dendritic cells and macrophages) overexpressed (but not the B7-1 ligand). B7-2 overexpression may be expected to be associated with increased T cell proliferation via IL-2 production</td>
<td>1998&lt;sup&gt;84&lt;/sup&gt;</td>
</tr>
<tr>
<td>---</td>
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<td>---</td>
</tr>
<tr>
<td><strong>Eosinophils and mast cells with IL-5 in acute disease</strong></td>
<td>Serum levels of both IL-5 (produced by Th2 cells, monocytes and mast cells) and eosinophils were increased in MMP. Mast cell and eosinophils were increased in the substantia propria in MMP. Mast cells released profibrogenic cytokines in pulmonary and hepatic fibrosis and may be important in MMP scarring.</td>
<td>Letko E 2002&lt;sup&gt;86&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Cytokine expression IFNγ, TGFβ, TNFα, PDGF</strong></td>
<td>Both diseased and normal control conjunctiva expressed IL-2, IFNγ (Th1 signature), TGFβ, TNFα, PDGF and bFGF as well as a proliferating cell marker. No IL-4 was expressed (Th2 signature). TGFβ and proliferating cells were overexpressed in the acute group and IL-2, bFGF and PDGF (fibrogenic cytokines) were more expressed in the subacute disease group.</td>
<td>Bernauer W 1993&lt;sup&gt;102&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>TGFβ1&amp; β3 increased</strong></td>
<td>TGFβ1&amp; β3 increased in the stroma in association with fibroblasts and macrophages in acute MMP (not chronic), PDGF and FGF undetectable</td>
<td>Elder MJ 1997&lt;sup&gt;87&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>HSP47, TGFβ and collagen I and III expression in conjunctiva and role of TGFβ 1 induction of HSP47 and collagen I by fibroblasts</strong></td>
<td>Collagen I &amp; III increased in OMMP conjunctival stromal tissue, and HSP47 and TGFβ increased in MMP fibroblasts. TGFβ1 induced HSP47 and collagen I in OMMP fibroblasts. Increased expression of the latter may contribute to conjunctival scarring in OMMP.</td>
<td>Razzaque MS 2003&lt;sup&gt;90&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Serum TNFα and IL-6 in active MMP</strong></td>
<td>Serum levels for TNFα, but not IL-6, were elevated in ocular MMP patients (n=35) compared to controls consistent with expectations for a systemic autoimmune disease.</td>
<td>Lee SJ 1993&lt;sup&gt;103&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>TNFα</strong></td>
<td>TNFα is overexpressed in inflamed OMMP conjunctiva compared to treated OMMP but the latter still shows greater expression than control levels. TNFα treatment of normal fibroblasts stimulated increased migration, mmp-9 production, decreased timp-2 and timp-4 and upregulated CD40 and ICAM. CD40 may interact with T lymphocytes</td>
<td>Saw VP 2009&lt;sup&gt;88&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>stimulating proliferation and profibrogenic cytokine production.</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td><strong>IL-13</strong></td>
<td>IL-13 is overexpressed in ocular MMP conjunctiva (both inflamed and non-inflamed) compared to controls. IL-13 promotes pro-fibrotic effects on normal human conjunctival fibroblasts and also upregulates the expression of T cell costimulatory molecules (CD80, CD40 and CD154) on fibroblasts potentially promoting interaction with T cells to drive fibrosis.</td>
<td>Saw VP 2009&lt;sup&gt;89&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>MIF protein and mRNA</strong></td>
<td>Increased MIF was demonstrated by immunohistochemistry and qPCR in both MMP conjunctiva and fibroblasts. The expression of MIF was associated with increased macrophage numbers. MIF was also secreted by fibroblasts (shown by ELISA). IL-1, TNF and TGFβ1 induced MIF expression in fibroblasts.</td>
<td>Razzaque MS 2004&lt;sup&gt;91&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>CTGF in stroma and fibroblasts</strong></td>
<td>CTGF is a downstream profibrotic mediator of TGF-β1 and was overexpressed in both ocular MMP whole conjunctiva and in cultured fibroblasts.</td>
<td>Razzaque MS 2003&lt;sup&gt;93&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>m-CSF</strong></td>
<td>Macrophage colony stimulating factor (m-CSF) was overexpressed in ocular MMP conjunctiva, macrophages, fibroblasts and epithelial cells. m-CSF correlated with macrophage numbers and some macrophages expressed a proliferation marker. Fibroblast m-CSF expression was increased by treatment with IL-1α or TNFα.</td>
<td>Razzaque MS 2002&lt;sup&gt;92&lt;/sup&gt;</td>
</tr>
<tr>
<td>Row</td>
<td>Immunosuppressive agent</td>
<td>% of patients or eyes responding to treatment</td>
</tr>
<tr>
<td>-----</td>
<td>-------------------------</td>
<td>---------------------------------------------</td>
</tr>
</tbody>
</table>
| 1   | Nicotinamide and tetracycline  
Reiche L 1998<sup>135</sup> | 63% (n=8) extraocular and 2 ocular MMP | Small case series. Rarely used for ocular disease. I have no experience of this therapy |
| 2   | Dapsone  
Saw VP 2008<sup>36</sup> | 77% (n=114) ocular MMP | Dapsone has common side effects, most often anaemia (95% of all patients to a variable extent), malaise and skin rashes. However, it’s not an immunosuppressive drug, may be very effective, and can also be combined with immunosuppressive drugs for added effect. Although unfashionable I still use it for these reasons. Compared to cyclophosphamide and steroid it is not as effective for severe disease (see below) |
| 3   | Rogers RS 1982<sup>108</sup> | 83% (n=24) extraocular and ocular MMP |  |
| 4   | Tauber J 1991<sup>133</sup> | 45% (n=69) ocular MMP |  |
| 5   | Foster CS 1986<sup>46</sup> | 70% (n=20) ocular MMP |  |
| 6   | Sulphapyridine  
Saw VP 2008<sup>36</sup> | 65% (n=55) ocular MMP | Usually now given as sulfasalazine (in which sulphapyridine is an active agent) because sulphapyridine is no longer widely available. I use this as an alternative when dapsone causes anaemia, or is not tolerated |
| 7   | Elder MJ 1996<sup>277</sup> | 50% (n=20) ocular MMP |  |
| 8   | Sulfasalazine  
Doan S 2001<sup>126</sup> | 45% (n=9) ocular MMP |  |
| 9   | Azathioprine  
Saw VP 2008<sup>36</sup> | 71% (n=80) ocular MMP | Azathioprine has similar effects to mycophenolate but side effects are more common such that mycophenolate has largely replaced azathioprine, except as a second line agent for patients not tolerating mycophenolate |
| 10  | Tauber J 1991<sup>133</sup> | 33% (n=11) ocular MMP |  |
| 11  | Ciclosporin  
Neumann R 1991<sup>132</sup>  
Foster CS 1992<sup>129</sup> | 2/22 patients (9%) ocular MMP | Calcineurin inhibitors have generally had poor success in ocular MMP and cannot be recommended on current evidence |
| 12  | Tacrolimus  
Letko E 2004<sup>130</sup> | 33% (n=6) ocular MMP |  |
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<tr>
<th></th>
<th><strong>Oral corticosteroids</strong>&lt;sup&gt;5&lt;/sup&gt;</th>
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<tbody>
<tr>
<td>13</td>
<td>Hardy KM 1971&lt;sup&gt;59&lt;/sup&gt;</td>
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<td></td>
<td>Mondino BJ 1979&lt;sup&gt;60&lt;/sup&gt;</td>
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<td></td>
<td>Foster CS 1986&lt;sup&gt;61&lt;/sup&gt;</td>
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<td></td>
<td></td>
<td>65% (n=23)&lt;sup&gt;6&lt;/sup&gt; ocular MMP</td>
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<td></td>
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<td>Doses above 0.5 mg/kg (about ≥ 40 mg) are usually needed to gain control of the disease and relapse is usual when doses are reduced. Side effects are unacceptable at the doses need as monotherapy Long term lower dose therapy has not been evaluated, but has not been needed in my patients having alternative steroid sparing immunosuppressive therapy. High doses (1-2mg/kg) are used for short term control in severe disease while the steroid sparing drug takes effect. I use a 3 months tapering course, usually in combination with cyclophosphamide</td>
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<tr>
<th></th>
<th><strong>Cyclophosphamide and steroids</strong>&lt;sup&gt;14&lt;/sup&gt;</th>
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<tr>
<td>14</td>
<td>Thorne JE 2008&lt;sup&gt;154&lt;/sup&gt;</td>
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<td></td>
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<td>91% (n=44) ocular MMP</td>
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<td></td>
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<td>This study used high dose (2mg/kg), prolonged (12-18 months) cyclophosphamide for most patients (68/78 or 87%). At 12 months 58/70 (83%) had complete success. However, serious complication rates, including malignancies (1 bladder cancers, 2 leukaemias and 6 other tumours), and 2 pneumonias, were relatively common and probably relate to a combination of the high doses used and the prolonged treatment period of 12-18 months which exceed safe doses (see Row 19)</td>
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<tr>
<th></th>
<th><strong>Elder MJ 1995&lt;sup&gt;128&lt;/sup&gt;</strong></th>
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<tr>
<td>15</td>
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<td>15/19 eyes (79%) ocular MMP</td>
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<tr>
<td></td>
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<td>2.0 mg/kg for up to 8 months</td>
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<tr>
<th></th>
<th><strong>Foster CS 1996&lt;sup&gt;90&lt;/sup&gt;</strong></th>
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<tr>
<td>16</td>
<td></td>
<td>100% (n=12) ocular MMP</td>
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<tr>
<td></td>
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<td>2.0 mg/kg</td>
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<tr>
<th></th>
<th><strong>Saw VP 2008&lt;sup&gt;50&lt;/sup&gt;</strong></th>
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<tbody>
<tr>
<td>17</td>
<td></td>
<td>90% (n=73)&lt;sup&gt;a&lt;/sup&gt; ocular MMP</td>
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<td></td>
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<td>1.0-1.5 mg /kg for up to 12 months</td>
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<tr>
<th></th>
<th><strong>Tauber J 1991&lt;sup&gt;133&lt;/sup&gt;</strong></th>
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<tr>
<td>18</td>
<td></td>
<td>92% (n=35) ocular MMP</td>
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<tr>
<td></td>
<td></td>
<td>2.0 mg/kg for a mean of 13 months (1-65 months)</td>
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### Additional notes on cyclophosphamide use:

- **Cyclophosphamide monotherapy.** Cyclophosphamide is typically used for patients with severe inflammation in ocular MMP, and is recommended by all authors as the drug of choice for this group of patients. It is usually combined with oral steroids for rapid disease control while the cyclophosphamide takes effect over 2-4 months. As a result, it is difficult to establish the effect of cyclophosphamide used as a single agent: in two of these studies 14/73<sup>36</sup> and 2/25<sup>133</sup> had cyclophosphamide without steroid. I use cyclophosphamide as single agent when patients have not responded adequately to other therapies, but do not have severe enough disease to risk oral prednisolone.

- **Cyclophosphamide dose and malignancy.** The dose of cyclophosphamide is given as between 1.0-2.0 mg/kg in the different studies and reviews summarized in rows 14-18. However, cyclophosphamide has a wide range of serious side effects listed in the studies below. In the last decade a maximum safe dose below which bladder cancer, and acute myeloid leukaemia are unlikely to occur, have been established in patients treated with cyclophosphamide for polyangiitis with granulomatosis. The probable cumulative safe doses for bladder cancer are 25gms, or 12 months treatment<sup>136</sup> and up to 36 gms, to limit the risk of both bladder cancer and acute myeloid leukaemia<sup>137</sup>.

- **How to use cyclophosphamide.** Cyclophosphamide tablets in the UK are available in 50mg or 100 mg and doses other than multiples of these can only be given by alternating therapy e.g. 50 mg one day alternating with 100 mg the next day to give a mean daily dose of 75 mg. I now use 1.0-1.5mg/kg<sup>36</sup> to restrict cumulative doses to the proposed safe limits. For most males I start with 100 mg per day for up to 12 months (a 36 gm cumulative dose for a 70kg person at 1.5 mg/kg). For women of 50kg I use 1 mg/kg for 12 months (an 18 gm cumulative dose over 12 months) but adjust the dose, and length of therapy, to get control of disease. Most patients will need 8 months of therapy to give them a period of one or two months of controlled inflammation and some will need more. This is less than the dose of 2.0mg/kg quoted in all the other studies, including our early publication<sup>128</sup>, to optimize safety. After the cumulative safe dose is reached I usually switch to a sulfa, before stopping cyclophosphamide, and add an alternative myelosuppressive if needed after discontinuing cyclophosphamide.
• **Intravenous cyclophosphamide.** The effects of this route have been rarely reported in ocular MMP case series: it is probably safer than daily oral therapy and may be as effective. However, for polyangiitis with granulomatosis a randomized trial of daily oral dosing versus intravenous cyclophosphamide (the CYCLOPS trial) showed that oral dosing was more effective in the long term. I have no experience of this modality of administration.

<table>
<thead>
<tr>
<th>No.</th>
<th>Medication</th>
<th>Study Details</th>
<th>Effects and Dosage</th>
<th>Other Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>Methotrexate</td>
<td>McCluskey P 2004</td>
<td>83% (n=12) ocular MMP</td>
<td>This is the only study describing methotrexate in detail. Other large studies have not used the drug (0/78) or only in small numbers (4/115), or the effects is not clear from the publication. I have little experience with this drug and use it as a third line agent when patients cannot tolerate, or don’t respond to, dapsone/suphasalazine, azathioprine, mycophenolate, and when cyclophosphamide cannot be used.</td>
</tr>
<tr>
<td>21</td>
<td>Mycophenolate</td>
<td>Saw VP 2008</td>
<td>82% (n=46) ocular MMP</td>
<td>Mycophenolate is one of the best tolerated and safest immunosuppressive drugs to use and is my first line recommendation for moderately severe disease.</td>
</tr>
<tr>
<td>22</td>
<td>IVIG</td>
<td>Zundel J 2001</td>
<td>9/10 eyes (90%) ocular MMP</td>
<td>This is a very expensive therapy and a scarce resource, but is approved for use in mucous membrane pemphigoid in the UK and accessible through local IVIG committees. I use this for patients who have failed other therapies. Supplementary Appendix 5 is a protocol for use in ocular MMP with acknowledgement to Foster CS, who has provided the evidence for its use in ocular MMP.</td>
</tr>
<tr>
<td>23</td>
<td>Infliximab, Etanercept</td>
<td>John H 2007, Prey S 2007, Sacher C 2002, Canizares MJ 2006, Heffernan MP</td>
<td>100% (n=7) ocular and extraocular MMP</td>
<td>These TNFα inhibitors have been little used in small case series. They probably deserve further study. I have no experience in their use for ocular MMP.</td>
</tr>
<tr>
<td>24</td>
<td>Rituximab</td>
<td>Le Roux-Villet C 2011, Taverna JA 2007, Ross AH 2009, Schumann T 2009, Lourari S 2010, Rubsam A 2015, Maley A 2016</td>
<td>80% (51/64) ocular and extraocular MMP although in addition to the 13 reported failures 3 patients in one series only had partial control and another 5 did not respond after late relapse.</td>
<td>Rituximab in these studies has been most often given using the rheumatology protocol in which one cycle is a 1gram infusion repeated 14 days later. All these patients have failed conventional therapy first. Repeat cycles given after 4 months or later have been effective in most patients not responding to one cycle. The effect of the drug in early cases, the relapse rate, safety and efficacy of adjunctive immunosuppressives, and long term management protocols have not been established. However, this is currently the most widely used rescue therapy for patients having failure to control inflammation by conventional therapy. I use this in patients in whom cyclophosphamide has failed or is contraindicated.</td>
</tr>
<tr>
<td>25</td>
<td>IVIG and Rituximab</td>
<td>Foster CS 2010</td>
<td>100% (n=6) ocular MMP</td>
<td>Very high doses of Rituximab were used in this study. I have no experience of combined therapy however I have recommended IVIG in Rituximab failures if two cycles of Rituximab 4 months apart have been ineffective 2 months after a failure to respond to Rituximab; at this stage there should be some residual effect from Rituximab.</td>
</tr>
</tbody>
</table>
27 Daclizumab
Papaliodis GN 2003
100% (n=1) ocular MMP
One case report only. I have no experience with this

28 Cyclophosphamide and prednisolone vs. oral prednisolone
Foster CS 1986
(n=24) 12/12 responded to cyclophosphamide and oral prednisolone versus 5/12 to oral prednisolone.
Confirms use of a steroid sparing drug with prednisolone versus prednisolone alone. No progression of disease in cyclophosphamide group versus 7/12 progressing in prednisolone group.

29 Dapsone versus cyclophosphamide
Foster CS 1986
(n=40) 14/20 responded to dapsone versus 20/20 to cyclophosphamide
Confirms the use of cyclophosphamide rather than dapsone for more severe disease.

30 Combination therapy with drugs other than prednisolone and cyclophosphamide

31 Tauber J 1991
Tauber J 1991: 105 patients. Combination therapy was used in 24% of patients at the end of follow up including dapsone +/- prednisolone +/- a myelosuppressive OR a myelosuppressive + prednisolone.
This influential paper was published over 2 decades ago and was the first to describe the effects of treatment of ocular MMP with most non-biological immunomodulators used today, in a large cohort of patients. The breakdown of outcomes of different therapies and combinations are less clear. However, combination therapy was used in 24% of patients and a recommendation is made for this in the paper.

32 Miserocchi E 2002
61 patients
22% requiring monotherapy, 21% requiring combinations of 3 or more drugs, and 10% 4 drugs

33 Saw VP 2008
Williams GP 2011
The same approach described in a different cohort of 54 patients.
388 treatment episodes in 115 patients: 198 single agent treatment episodes (51%) with success or partial success in 73%. 93 treatment episodes with a myelosuppressive (cyclophosphamide, azathioprine, mycophenolate, methotrexate) OR a sulfa (dapsone, sulphapyridine) + prednisolone having 87% success or partial success. 47 treatment episodes with a myelosuppressive + a sulfa having 75% success or partial success. 31 treatment episodes with a myelosuppressive (cyclophosphamide, azathioprine, mycophenolate, methotrexate) + a sulfa (dapsone, sulphapyridine) + prednisolone having an 87% success or partial success rate. The paper includes the data on the outcomes of all 115 patients as supplementary data.

34 Additional notes on combination therapy
• Saw VP 2008 is the most comprehensive study on treatment methodology for conventional (non-biological immunomodulatory) therapies. The paper includes outcomes of single agent therapy, and multiple agent combinations with rates of side effects. Multiple agent therapy resulted in improved response rates compared to single therapy, without an identifiable increase in complication rates over those for monotherapy. A “step up and step down” approach is described with the choice of initial therapy related to the severity of disease. Additional drugs are added to, as combination therapy, or substituted as monotherapy, when necessary to gain disease control or to avoid side effects. Cyclophosphamide is reserved for the more severe cases, less toxic drugs for others, and combinations of sulfas and myelosuppressives when disease control on monotherapy is inadequate. Prednisolone, used in short courses, was added to any of the other drugs but most widely in combination with cyclophosphamide.
• Guidelines for cyclophosphamide therapy are given in Row 19. Dapsone or sulphapyridine was either added to myelosuppressives, when reducing therapy
• This regimen is what is currently used in the Moorfields Eye Hospital Clinics. This approach has been adopted in UK specialist centres and is also what has been described also been described in the Foster publications summarized above. However, three review articles have not mentioned the use of combined therapy, excepting the use of prednisolone as adjunctive therapy, and have not discussed the value of step up and step down therapy, using drug combinations, to maintain therapy during treatment changes and to minimize risk, for ocular MMP. The approach used by Thorne JE et al. 2008 using cyclophosphamide as monotherapy after a course of adjunctive prednisolone is effective but probably less safe (see Rows 14,19)

Supplementary notes:

a results for success (fully controlled inflammation) and qualified success (partial control of inflammation) of 388 individual treatment episodes in 115 patients
b combined results of three case series
c combined results of two case series
d combined results of five case series
e combined result of 3 case series and 4 case reports