C1 inhibitor deficiency: consensus document

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Introduction

This document was commissioned by the Primary Immunodeficiency Association (PIA). It represents a consensus from patients, experts and the literature on the diagnosis, therapy and management of C1 inhibitor (C1 INH) deficiency.

For the purpose of this document C1 INH deficiency will include both genetic [types I and II hereditary angi-oedema (HAE)] and acquired [acquired C1 inhibitor deficiency (formerly acquired angi-oedema, AAE)] forms of the disease. It should be noted that this is a rare disorder and much of the literature is based on case studies or small series. The syndrome of type III HAE is referred to where appropriate, but is not part of the spectrum of C1 INH deficiency and as such is not covered in depth in this document. The levels of evidence used are listed in Table 1.

Background

C1 esterase inhibitor deficiency [hereditary or acquired (HAE/AAE)] is characterized by the occurrence of subcutaneous and submucosal swellings in any part of the skin and the respiratory and gastrointestinal tracts. In the hereditary form, symptoms usually appear early in life and are normally accompanied by a family history. Although scattered reports of this disease can be traced back to the last century, hereditary angi-oedema reached its own identity in 1963 (for reviews see Cicardi et al. [1] and Fay and Abinun [2]).
Genetics and prevalence

The disease is inherited in an autosomal dominant manner. The spontaneous mutation rate is about 25% and more than 100 different C1 inhibitor gene mutations have been described [3]. The prevalence of the disease has been estimated at 1/50 000, with no reported bias in different ethnic groups.

While it is unusual to find the disease without symptoms, there is an extreme variability in their frequency and severity [4]. There seems to be little, if any, correlation between symptoms and type of genetic defect with patients from the same family, and therefore sharing the same mutation, showing wide differences in phenotype [4].

In HAE type I (up to 85% of all patients), there is a deficiency in the amount of C1 INH protein present in the plasma. This is the result of only one gene functioning. However, plasma values are usually 5–30% of normal rather than the 50% value that might be expected [5]. Increased catabolism of C1 INH, even in asymptomatic patients, and possibly decreased production, are likely factors [3,5]. There is also some evidence that certain amino acid substitutions found in type I HAE affect the intracellular transport of C1 INH and result in a marked reduction or the total impairment of protein secretion [3].

In HAE type II, the circulating C1 INH concentration is normal or high but not fully functional. In vitro studies show that C1 INH production in type II HAE is normal in contrast to the findings in patients with type I disease [5]. High plasma concentrations of dysfunctional C1 INH are found because the mutant protein is secreted normally and it is unable to form complexes with proteases, which increases its half-life in the circulation. Dysfunctional proteins often result from substitutions at the reactive site loop Asp446, but may also result from changes at several positions outside the reactive site loop.

HAE type III has been described by Bork et al. [6]. In this paper, cases with typical clinical features of C1 INH deficiency were described with normal C1 INH level and function and a normal C4. These cases were all female and seemed to have a dominant mode of inheritance.

AAE is said to affect a tenth as many patients as HAE, although this may be an underestimate. AAE presents in older patients, has no family history and is associated with lymphoproliferative disease or, less commonly, autoimmunity [7,8].

Immunology

C1 INH is the main regulator of the early activation steps of the classical complement pathway. This protein is produced mainly in the liver, but also by activated monocytes and other cell types [9]. C1 INH also regulates the activation of kallikrein, plasmin in the fibrinolytic pathway, the activation of factor XI in the coagulation cascade and activated factor Xlla. In the presence of C1 INH deficiency the classical complement pathway can be inappropriately or excessively activated. Immune complexes trigger the activation of the first component C1 to C1 esterase. C1 esterase then acts with its natural substrates C4 and C2 to form the complex C4b2a. Formation of this new complex (and associated C3 activation) leads to the production of anaphylactic, chemotactic and vasoactive peptides (C2b, C3a, C5a). C1 INH protein blocks both the spontaneous activation of C1 and the formation of activated C1, therefore not allowing the C2,4 complex to be created.

In the kinin releasing system, C1 INH regulates conversion of prekallikrein to kallikrein. C1 INH deficiency results in an increase in kallikrein, which in turn increases bradykinin production. Inhibitory effects of C1 INH in factor XIa, factor Xla and plasmin have also been described. The end result is increased vascular permeability and massive local uncontrolled oedema. While there is some debate as to the exact component that contributes to the angio-oedema, there is accumulating evidence to support the involvement of bradykinin [4,10–12].

Diagnosis

Clinical

A diagnosis of C1 INH inhibitor deficiency is suggested by a history of recurrent attacks of angio-oedema and of abdominal pain (see Table 2). Symptoms include recurrent circumcribed, non-pruritic, non-pitting oedema. Peripheral pain is not usually a feature, unless swelling occurs on pressure bearing areas or where subcutaneous tissue is limited. Oedema can affect virtually any part of the integument, but is more common in the extremities [13]. Episodes of swelling may also involve the upper respiratory tract, including the tongue, pharynx and larynx. This contributed to the 15–33% mortality from the disease reported previously in the literature [14]. Abdominal pain, nausea and vomiting are the dominant symptoms in approximately 25% of all patients, and are the result of constriction by intestinal wall and mesenteric oedema [15]. Urticaria is not a feature of C1 INH deficiency. However, prodromal erythema has been reported in up to 25% of patients which may be mistaken for urticaria [16,17].

Classically, the oedema and swelling develop gradually over several hours, increasing slowly for 12–36 h, and then subside after 2–5 days. However, patients may experience
abdominal attacks with a very sudden and severe onset of pain and no visible oedema. Attacks of severe swelling can occur in some patients on a weekly basis and in others happen only once or twice a year.

Angio-oedema can be precipitated by minor trauma to the tissue, such as dental work (said to be a cause in up to 50% of all cases) [18,19], by certain drugs such as oestrogen [20] or angiotensin converting enzyme inhibitors, by emotional stress or by infection [21].

Acute attacks of abdominal pain can mimic surgical emergencies and, before a diagnosis of HAE is established, patients frequently undergo unnecessary appendicectomy or exploratory laparotomy. Equally, after diagnosis, there is always the concern that true abdominal emergencies will not have surgery performed in good time [4]. Barium studies, carried out during an acute attack, show massive submucosal oedema, spiculation and fold thickening or effacement [22]. The gastrointestinal involvement appears to be segmental and transient with reversion to normal by several days after an attack. In a report of an endoscopy carried out during an acute attack of C1 INH deficiency the gastric mucosa was described as diffuse reddish and oedematous and the mucosal surface in involved areas bulged remarkably, mimicking a submucosal tumour [23]. Histological examination of the bulging area merely showed moderate inflammatory cell infiltration of the lamina propria [23]. These findings are relatively non-specific and response to treatment with C1 INH concentrate may be the only way to differentiate a surgical condition from an acute attack of C1 INH deficiency [4].

### Laboratory

Laboratory tests should be performed in an accredited laboratory registered with a suitable quality assurance scheme (e.g. UK National External Quality Assessment Scheme). Serum C4 level is a good screening test for C1 INH deficiency as serum C4 is invariably low in untreated HAE (C4 < 30% of mean normal level) [24]. It has been shown that for untreated C1 INH deficiency low C4 has 100% sensitivity, 100% negative predictive value and is thus an effective screening test [24]. All patients who are suspected of having C1 INH deficiency should have a C4 level measured. If C4 is normal it is not usually necessary to proceed to C1 INH analysis [24]. If the C4 level is low then C1 INH level and function should be assessed.

The diagnosis of type I HAE (85% of cases) is by demonstrating low amounts of C1 inhibitor protein, as assessed by immunochemistry. If C1 inhibitor value appears normal or raised (and C4 is low), a test of C1 inhibitor function should be carried out [18,25]. An absence of function suggests a type II defect. All such tests should be carried out on a fresh (or freshly frozen) serum sample, i.e. one less than 4 h old.

If C1 INH function or/and level are low and C4 is low then a repeat sample should be obtained to confirm the findings. The low prevalence of the condition means that false positives are common [24]. All testing should be undertaken off treatment, including the administration of C1 INH concentrate or fresh frozen plasma, to allow reversion to untreated levels. Ideally this should be for more than a week but longer if borderline levels are obtained.

Interpretation in very young children is difficult, owing to a paucity of data regarding reference ranges in children. C4 is not a reliable indicator in the very young as, again, the reference range is extended downward with respect to the adult reference range [26]. Data suggest that C1 INH is reduced by 30–50% in normal neonates (cord blood analysis), both antigenically and functionally [27]. In children under 1 year of age a low C1 INH (less than 30% mean adult level) confirms the diagnosis of HAE. However, the diagnosis cannot be excluded in a child under 1 year of age, even if the C1 INH level and function are normal. In this case, investigations should be repeated when the child is over 1 year.

There is evidence that pitfalls in the diagnosis are common, with 11 of 42 cases reviewed recently found to have a questionable diagnosis [28]. Established or transferring cases should be reviewed for validity of the diagnosis. In the presence of a low C1 INH level or function but a normal C4 the diagnosis of HAE must be questioned. We would advise that, in these circumstances, C1 INH be rechecked by a different method (evidence level 4). Currently, genetic tests are not indicated routinely; however, under certain circumstances a genetic test may be of use, if available. In cases where the diagnosis is established, C4 and levels of C1 INH and function may be useful to monitor treatment effect.

### Management

**Primary prevention**

Management of patients with C1 INH inhibitor deficiency should cover their long-term, short-term and acute needs. It is important in the general management of these patients to search for potentially treatable triggers of attacks and deal with them.

<table>
<thead>
<tr>
<th>Table 2. Diagnostic features which should prompt investigations for C1 inhibitor deficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angio-oedema</td>
</tr>
<tr>
<td>Recurrent</td>
</tr>
<tr>
<td>&gt;24 h</td>
</tr>
<tr>
<td>Non-pruritic</td>
</tr>
<tr>
<td>Non-responsive to antihistamines</td>
</tr>
<tr>
<td>Serpiginous rash</td>
</tr>
<tr>
<td>No urticaria</td>
</tr>
<tr>
<td>Unexplained abdominal pain</td>
</tr>
<tr>
<td>Recurrent</td>
</tr>
<tr>
<td>‘Colicky’</td>
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<tr>
<td>Family history</td>
</tr>
<tr>
<td>Low C4</td>
</tr>
</tbody>
</table>
Infected teeth and other foci of infection, which may activate complement, should be sought and treated [29,30]. Eradication of Helicobacter pylori may be beneficial [31,32] (evidence level 3). Patients and their general practitioners (GPs) should be advised that infections should be treated promptly.

Advice on use of contraceptives and hormone replacement therapy should emphasize avoidance of oestrogen (see below, ‘Special situation-contraception’). Angiotensin-converting enzyme (ACE) inhibitors need to be avoided because of their effects on the kallikrein–bradykinin pathway [33]. Both HAE and AAE may be manifest for the first time after treatment with ACE inhibitors or oestrogens [34,35] (evidence level 3). Angiotensin-II receptor antagonists may also induce angio-oedema in normal patients [36,37], although the majority of patients with ACE-inhibitor-induced angio-oedema tolerate angiotensin-II receptor antagonists [38]. Angiotensin-II receptor antagonists may be used with caution in patients with C1 INH deficiency. Beta-blockers or diuretics should be considered first-line treatment for hypertension in patients with C1 INH deficiency. There is no evidence for or against the use of methyl dopa to control hypertension in this group of patients.

Attacks are likely to become more frequent at times of physiological or psychological stress, so it may be sufficient to use prophylactic drugs during such periods only, thus minimizing adverse effects. Nevertheless, there will be a group of patients who will require intermittent or continuous, long-term prophylaxis.

The threshold for treatment should be a joint decision between clinician and patient. It should include an assessment of the severity, frequency and life-threatening nature of the attacks. This, in combination with the patient’s circumstances, should allow the development of an appropriate treatment plan.

### Long-term prophylaxis

The regimen for each affected individual should be guided by the severity of the disease. Frequent attacks of peripheral angio-oedema (extremities, trunk), although unpleasant and annoying, are not dangerous and may not require (contingent upon the patient’s judgement) long-term prophylaxis. However, prophylactic administration of antifibrinolytic agents (tranexamic acid [39] or epsilon-aminocaproic acid (EACA; not licensed in the UK) [40]) and/or synthetic attenuated androgens (danazol [41–43] or stanozolol [43–46]), has proved useful in reducing the frequency or severity of attacks (evidence level 2/3). Other androgens (methyltestosterone [47], fluoxymesterone [48] and oxymetholone [48,49]), can be used in adult males (evidence level 3). Non-17 alpha alkylated derivatives, such as nandrolone, appear to be ineffective and should not be used [43].

A graded approach to the level of treatment can be tailored to the individual with minor peripheral episodes. Consideration could be given to a course of tranexamic acid before attenuated androgens. Maintenance treatment should be considered in any patient who has had more than one episode of severe abdominal pain in one year or any head or neck swellings, frequent peripheral or genital swellings or a requirement for concentrate more than once a year. Fatal episodes have occurred in patients who previously have had only mild or benign attacks [50] (evidence level 2).

### Antifibrinolytic agents

Antifibrinolytic agents inhibit plasminogen activation with consequent ‘sparing’ of C1 INH usage. They decrease the number and the severity of attacks [19], but are not as effective in this as the attenuated androgens [40] (evidence level 2). Their side effects include nausea, vertigo, diarrhoea, postural hypotension, fatigue and muscle cramps with an increase in muscle enzymes concentrations [1,39,40,51–53] (evidence level 2/3), and theoretical concerns about thrombus formation and thrombotic episodes [18]. However, recent reports have suggested that these side effects are less common than thought previously; long-term use in menorrhagia has shown no evidence for increased thrombus formation [53]. The finding of tumours of the retina and liver in experimental animals after long-term use of tranexamic acid [18] has limited its use in the United States [15], but not in Europe [54,55]. Although a teratogenic effect of EACA has been postulated in the period of embryonic growth and development [18,56] it is being used in the United States [57], it has been used in children [58] and, surprisingly, has been recommended during pregnancy [59].

A starting dose of 1–1.5 g of tranexamic acid up to two to three times a day [60] should be used depending on disease severity, reducing to 0.5 g once or twice a day as the attacks remit. In children, the dosage will need to be adjusted (see Table 3). Diarrhoea may be a limiting side effect. Patients should be warned of this possibility and, if necessary, the dose titrated against side effects (evidence level 4).

Although there is no evidence of teratogenicity from animal studies, we recommend avoiding the use of tranexamic acid in pregnancy, if possible. The British National Formulary (BNF) indicates that regular eye examinations and liver function tests (LFT) should be performed, while recognizing that the evidence base for this is minimal. We suggest that fundoscopy should be performed annually, with referral if symptoms occur, and LFT performed every 6 months.

### Attenuated androgens

Attenuated androgens increase the biosynthesis of many proteins, including the hepatic production of C1 inhibitor protein [18]. Danazol, stanozolol and oxandrolone are most commonly used. Their side effects, which are dose-dependent, include weight gain, virilization, muscle pains and cramps, headaches, depression, fatigue, nausea, consti-
pation, menstrual irregularities and liver function derangement [46,61,62] (evidence level 3). Decreased growth rate in children [63–65] is the main contraindication for their use in this age group. Androgens can cause masculinization of the female fetus [66,67] and thus are contraindicated during pregnancy. Androgens, particularly the 17-alpha alkylated androgens, may have hepatic side effects, including cholestatic jaundice [68], peliosis hepatitis [69] and hepatocellular adenoma [70–73]. The observed cases of hepatocellular adenomas developing in patients with C1 INH deficiency on long-term prophylaxis with danazol have caused particular concern [74] (evidence level 3). A dose of danazol 200 mg once or twice a day will usually suffice in adults, preventing attacks in 80% of cases [7] (evidence level 2). Because of the wide variations between individuals with this condition the dosage must be titrated to individual need and up to 400 mg twice a day may be required. Conversely, once symptom control is established, many patients remain well on doses as low as 100 mg thrice weekly. Stanozolol at a dose of up to 5 mg once or twice daily can be used where available [46]. To facilitate more accurate titration of dosage a 2 mg tablet has been introduced. Stanozolol is available in the United Kingdom

### Table 3. Treatment summary

The regimen for each affected individual should be guided by the severity of their disease and thus titred to individual need. The following is a guide to the dosage and summarizes the advice given in the text.

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Therapy</th>
<th>Dosage (adult)</th>
<th>Dosage (children)</th>
<th>Monitoring tests</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Long-term prophylaxis</strong></td>
<td>Attenuated androgens</td>
<td>Danazol 200 mg once or twice per day; up to 400 mg/day in &lt;20% of cases</td>
<td>[Only if indicated (very rare), see text]</td>
<td>Six-monthly: liver function tests</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Stanozolol up to 5 mg once or twice per day</td>
<td>Danazol 100–200 mg/day (use lowest effective maintenance dose, consider alternate-day or 2× weekly regimen)</td>
<td>Annual: lipid profile</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Oxandrolone 2·5–20 mg divided dose 2–4 times per day (use lowest effective maintenance dose, consider alternate-day or 2× weekly regimen)</td>
<td></td>
<td>Biennial: hepatic ultrasound (annual after 10 years’ treatment)</td>
</tr>
<tr>
<td></td>
<td>Tranexamic acid</td>
<td>Starting dose 1–1·5 g 2–3 times per day, reducing to 0·5 g once or twice per day</td>
<td>1–2 g per day; dosage depends on age and size; general guide is 50 mg/kg/day (use lowest effective maintenance dose, consider alternate-day or 2× weekly regimen)</td>
<td>Six-monthly: liver function tests</td>
</tr>
<tr>
<td><strong>Short-term prophylaxis</strong> (e.g. for dental work)</td>
<td>C1 inhibitor concentrate</td>
<td>500–1500 U up to 24 h prior to procedure</td>
<td>&lt;10 years old 500 U, &gt;10 years old 1000 U up to 24 h prior to procedure</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Attenuated androgens</td>
<td>Danazol 100–600 mg/day for 48 h before and after procedure</td>
<td>Danazol 300 mg/day for 48 h before and after procedure</td>
<td></td>
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<tr>
<td></td>
<td>Tranexamic acid</td>
<td>Stanozolol 2–6 mg/day for 48 h before and after procedure</td>
<td></td>
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<tr>
<td></td>
<td>Fresh frozen plasma</td>
<td>1 g given four times daily for 48 h before and after procedure</td>
<td>500 mg given four times daily for 48 h before and after procedure</td>
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</tr>
<tr>
<td><strong>Emergency care for acute attacks</strong></td>
<td>C1 inhibitor concentrate</td>
<td>500–1500 U; additional infusion and reassessment if symptoms persist for &gt;2 h</td>
<td>&lt;10 years old 500 U, &gt;10 years old 1000 U</td>
<td>Baseline: liver function tests, hepatitis virology</td>
</tr>
<tr>
<td></td>
<td>Attenuated androgens</td>
<td>Danazol up to 1 g/day</td>
<td></td>
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<tr>
<td></td>
<td>Tranexamic acid</td>
<td>Stanozolol up to 16 mg/day</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fresh frozen plasma</td>
<td>1 g given four times daily for 48 h</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>2 units (only for use where Cl INH concentrate not available)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pain relief</strong></td>
<td>As appropriate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pregnancy</strong></td>
<td>Attenuated androgens</td>
<td>Contraindicated</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tranexamic acid</td>
<td>May be used with caution</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>C1 inhibitor concentrate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Emergency care as above. Severe cases may require regular replacement</td>
<td></td>
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</tbody>
</table>
only by importation and on a ‘named patient’ basis. The recommended adult dose for oxandrolone is 2·5 mg to 20 mg given in two to four divided doses [75]. Again, the doses of these should be titrated according to individual need. In some cases combined therapy, e.g. attenuated androgens plus tranexamic acid, may be beneficial.

Some male and many female patients experience troubling or unacceptable side effects on their prescribed dose of attenuated androgens. It is important to explain the advantages and disadvantages of the treatment regimen, to discuss fully possible side effects with the patient and to monitor regularly the acceptability of such side effects.

**Long-term C1 INH prophylaxis**

Long-term prophylaxis with C1 INH may be necessary in patients where tranexamic acid or steroids are not effective, not tolerated or contraindicated. This may include those with underlying thromboembolic disease or during pregnancy. Prior to recommending regular therapy, access to C1 inhibitor for acute attacks should be optimized, by home therapy training if necessary. In exceptional cases where this approach does not provide sufficient symptom control, regular C1 inhibitor infusions of 500–1000 U twice weekly may be required.

**Short-term prophylaxis**

Short-term prophylaxis for surgical procedures is the third arm of treatment in these patients. If surgery or dental work is to be carried out on a planned basis, an infusion of C1 inhibitor concentrate can be given up to 24 h before the procedure [25], or just prior depending on the individual circumstances [55,76]. It is impossible to predict the requirements of an individual patient in such a situation; body mass and previous requirements will be helpful indicators. In general, an infusion of 1000 U of concentrate should be sufficient for most dental work and most planned surgery for an adult patient but requirements may vary from 500 to 1500 U. A further dose may be required, particularly if there is postoperative infection.

Administration of antifibrinolytics or attenuated androgens, starting 5 days before the procedure and the following 2 days thereafter [54], is an alternative. There are no data on the relative efficacy of concentrate to attenuated androgens in this setting. Tranexamic acid has been used at a daily dose of 4 g (1 g four times daily) for adults [77,78] or 2 g (500 mg four times daily) for children [55], given 48 h before and after surgery. However, it seems that most authors prefer attenuated androgens, where concentrate is not used, even in children [18,54] at a dose of 100–600 mg/day for danazol or 2–6 mg/day for stanazolol, given 48 h before and after surgery [1,18,46,54]. See below, 'Dental care', for further information on dental care.

**Patient possession of C1 INH concentrate**

All C1 INH deficiency patients should be offered the opportunity for home possession of C1 INH, of a sufficient therapeutic dose to treat a laryngeal emergency, as 50–75% have a life-threatening attack at some time [1,79].

A UK audit has shown that home possession could reduce the number of avoidable adverse effects [80] (evidence level 2). In order to be effective good local links to accident and emergency and a care management plan are also essential.

**Home possession – patient-directed administration**

The management of patient-administered C1 INH concentrate is in need of standardization. Therefore, the attached recommendations (adapted from the TRIC Guidelines for Home Therapy and Home therapy for C1 INH deficiency, St Bartholomew’s Hospital) are put forward as example assessment guidelines to be instituted prior to home therapy being initiated (see Table 4).

Home therapy requires the issuing of concentrate and the training of participants of all eligible patients with C1 INH deficiency (see Table 5). It provides a quick, convenient and
probably a safe method of dealing with acute attacks of angioedema [80]. This is particularly valuable where access to emergency care is likely to be difficult through reasons of resource or geography.

However, there are also a number of important safety considerations. There has to be provision of refrigeration facilities for the storage of the product. Reassuringly, experience has shown that the product retains efficacy for many months under less than optimal storage conditions (e.g. 6 months at 25°C) (evidence level 4). Very recently a new formulation has undergone a series of room temperature storage tests and has shown good long-term stability [81]. Experience with C1 INH indicates that adverse reactions are very rare. Home therapy programmes with intravenous immunoglobulin have demonstrated that it is possible to train patients, with an ‘infusion partner’, to manage infusions and adverse events safely at home. Because C1 INH is likely to be required when the patient is unwell, 24-h emergency treatment at the local hospital must remain an option. Patients and carers should be encouraged to use this option where appropriate. There is evidence that self-possession reduces the time patients spend awaiting infusions [80] (evidence level 2).

Any such programme should be accompanied by appropriate information to be carried with the patient and advice as to strategies for resupply of concentrate.

Monitoring side effects of treatment

**Tranexamic acid**

The BNF recommends that patients who receive long-term tranexamic acid have a regular eye examination, but notes that this is based on unsatisfactory evidence [60]. The BNF further recommends regular checks of liver function (evidence level 4).

Use of tranexamic acid is contraindicated in active thromboembolic disease. Hence, if there is a personal or family history of thromboembolic disease, we suggest a thrombophilia screen should be performed before commencing treatment (evidence level 4).

**Attenuated androgens**

Liver function tests should be performed every 6 months; both tranexamic acid and attenuated androgens can cause abnormalities. Danazol and other attenuated androgens may affect lipid metabolism and thus confer an added risk of cardiovascular disease. Therefore, lipids should also be checked at presentation. Fasting lipids need testing only where initial screening is abnormal. Thereafter we recommend checking at 6 months and 1 year. When patients have no increase in their attenuated androgen dose, no weight or dietary change, both tranexamic acid and attenuated androgens can cause abnormalities. Danazol and other attenuated androgens may affect lipid metabolism and thus confer an added risk of cardiovascular disease. Therefore, lipids should also be checked at presentation. Fasting lipids need testing only where initial screening is abnormal. Thereafter we recommend checking at 6 months and 1 year. When patients have no increase in their attenuated androgen dose, no weight or dietary change, if lipids are stable after 12 months, further checking annually is sufficient (level of evidence 4).

**Hepatic ultrasound**

The report of hepatocellular adenomas developing in patients with C1 INH deficiency on long-term prophylaxis with danazol [74] (evidence level 3) has indicated that ultrasound screening may be useful. As yet there are no data to indicate the extent of the problem or the frequency of screening required.
Danazol and other 17-alpha-alkylated steroids are associated with increased risk of peliosis hepatis and hepatic adenoma [82]. We recommend that all patients taking regular or frequent courses of attenuated androgens should have a baseline ultrasound, which should be repeated every 2 years, or annually in patients who have been treated for more than 10 years. This recommendation is based on expert opinion (level 4) [74,83–85].

Emergency care

Treatment of acute attacks depends on their severity. Episodes of peripheral swelling only usually do not require treatment, but stanozol (up to 16 mg/day [86]) or danazol (up to 1 g/day) given early during an attack may shorten its duration. Involvement of the upper airway usually begins slowly but cases of progression within 20 min have been reported [50]; voice alteration and dysphagia indicate high risk of total airway obstruction. If there is any suspicion of airway involvement C1 INH concentrate should be given promptly. The dose requirement will vary between individuals, dependent on body mass and the seriousness of the condition. In a life-threatening situation we recommend 1000–1500 U. In other situations 500–1000 U is often sufficient. Administering C1 INH concentrate shortens the duration of attacks by about a third and also halves the time to the beginning of the relief of symptoms [25].

For acute attacks of abdominal oedema, pain relief should be given at an appropriate level. Non-steroidal anti-inflammatory drugs are useful in the treatment of abdominal pain. If the attack is severe, C1 INH concentrate should be infused at the same dose as above. Early intervention prevents avoidable pain and reduces disruption to the patient’s life. The patient should be observed closely until symptoms start to improve. The median time to the beginning of the relief of symptoms after concentrate infusion is 0.5–1.5 h, with complete resolution of symptoms after 24 h [25]. If symptoms persist at a high intensity 2 h after infusion, additional C1 INH concentrate should be given and alternative diagnoses should be considered.

C1 INH concentrate is available throughout Europe. It has been available since the early 1980s [87], and shown to be effective in case series and a controlled trial [76,88,89] (evidence level 2). If concentrate is not available then fresh frozen plasma (FFP) or solvent detergent-treated plasma may be given (evidence level 3), although this may worsen symptoms during the acute phase [15,18,56] because it contains high concentrations of complement components. A solvent/detergent-treated plasma (Octaplas, Octapharma AG, Vienna, Austria) has been evaluated for use in HAE, but there are few data regarding efficacy [90].

There are no randomized trials comparing plasma with C1 INH concentrate or with placebo. The risk of pathogen transmission may be increased if plasma is used [91,92]. Therefore, plasma is not an acceptable alternative where emergency treatment is foreseeable.

Adrenaline is used to treat angio-oedema and hypovolaemia associated with type 1 hypersensitivity. In the context of HAE there is little evidence that, relative to other treatments, it is efficacious.

Potential new therapies

New inhibitors of the fibrinolytic system, such as the kallikrein inhibitor DX88 (Dyax Corp., Cambridge, Massachusetts, USA) and the bradykinin B2 receptor inhibitor Icatibant (Jerini AG, Berlin, Germany), hold promise for use in the treatment of C1 INH deficiency [93] and trials are commencing. Recombinant C1 inhibitor (Pharming Group NV, Leiden, Netherlands) has been developed and trials should be available in the near future [94].

Special situations

In pregnancy and delivery

Treatment of the disease during pregnancy has special problems. Of published reports, some anecdotes report worsening of the disease [95] (evidence level 3), but few attribute premature labour or stillbirths to the disease [96,97]. In a series of 25 pregnancies in affected patients, only two had an increase in frequency of attacks, and none of these was related to the delivery itself [18] (evidence level 2). Ideally, all prophylactic drugs should be stopped during pregnancy and, if possible, before conception. Of particular note, attenuated androgens are contraindicated during pregnancy [98]. If prophylaxis is required, tranexamic acid may be used with caution. Although tranexamic acid crosses the placenta, there are no data to suggest that tranexamic acid is teratogenic. Further, there does not appear to be an increase of thromboembolic events [99] (evidence level 2). Severe attacks during pregnancy should be treated with concentrate as in the non-pregnant patient. Severe cases may require regular C1 INH replacement therapy.

There is little evidence that complications from C1 INH deficiency are common in vaginal delivery. The consequences of an attack during delivery are potentially serious. Individual patients should be discussed with the obstetrician. Consideration should be given to the obstetric risk (e.g. primigravida), C1 INH deficiency history and the patient’s own views. The safest obstetric approach would appear to be to administer a predelivery infusion of 500–1000 U C1 INH concentrate [100]. In a low-risk pregnancy without pretreatment with C1 INH concentrate we recommend that C1 INH should be available in the delivery suite. There may be local swelling of the vulva and infusion sites, but this would not be treated unless urethral obstruction was a problem [21]. If an operative delivery is undertaken, regional analgesia is to be preferred to endotracheal intubation in order to avoid laryn-
geal trauma [101] (level of evidence 3). In all situations the clinician should consider the postpartum period one of higher risk of acute attacks.

**Contraception**

Oestrogens should be avoided where possible. The use of combined oral contraceptives exacerbate symptoms in HAE patients [102,103] (evidence level 3). A recent study reported that over 60% of HAE types I and III patients have more frequent attacks on oestrogens [20] (evidence level 2). In general, progesterone-only pills such as levonorgestrel are preferred. Progesterone may have a mildly protective effect. No published data exists regarding the use and safety of intrauterine devices.

**Dental care**

Trauma can precipitate acute oedema in patients with C1 INH deficiency. For this reason dental work carries a risk of triggering an attack. Fatal laryngeal attacks have been reported following tooth extraction [88]. However, attacks are unpredictable. Extensive dental work may be carried out without complication and conversely minor work may sometimes precipitate an attack [104].

All patients should be warned of the increased risk of an attack in the 36 h following dental procedures and should have rapid access to C1 INH replacement in the event of an attack [105], irrespective of whether they have received prophylaxis. Recommendation for prophylaxis should take account of the proposed dental procedure and of previous reactions experienced by the patient.

Danazol, C1 INH concentrate and FFP have all been recommended for prophylaxis [88,104,106]. We believe that correction with C1 INH is to be preferred for more invasive dental procedures (e.g. extractions). It is more physiological than treatment with attenuated androgens and is more likely to reliably achieve normal levels of C1 INH. Furthermore, use of C1 INH overcomes any potential doubt regarding adherence with anabolic steroids (evidence level 4).

**Travel**

The following advice is taken from the Primary Immunodeficiency Association (PIA) advice for HAE patients when travelling in the United Kingdom and abroad. Further advice can be obtained via the PIA website [http://www.pia.org.uk]. The advice falls into two broad categories: general administrative advice and that related to emergency treatment.

**General advice**

Wear a Medic Alert bracelet. Obtain form E111 from your local post office if travelling in Europe. Arrange travel insurance that will cover HAE. Discuss the situation well in advance with your consultant for advice on medication and emergency treatment. A doctor’s letter will be required in order to take C1 INH through airport controls. Medications should be declared at the baggage checks and carried as hand luggage in a coolbag.

**Emergency advice**

Carry a consultant’s letter giving instructions about emergency treatment and a 24-h emergency advice telephone number (translated if travelling abroad). All HAE patients should have an emergency dose of C1 INH to keep with them when travelling away from their home base, as well as standard treatment.

**Children**

Attacks are seen during childhood in most patients [18,107]. Although the diagnosis is usually made in the second or third decade of life [18,108,109], it is well documented that between 50% and 75% of patients had their first attack by the age of 12 years. Data from the largest patient group studied (over 340 patients from 120 different kindreds) and followed over a period of more than 20 years [1,4,7,54,110] confirms that almost 40% had onset of their symptoms before the age of 5 years, and 75% before the age 15. Data from smaller studies on children only provide more striking evidence that most experienced their first symptoms in early childhood, before the age of 6 years [58,111]. Occasional patients will have their first symptoms even earlier, before the age of 1 year [27,110,112,113]. Attacks in children are usually not as frequent and/or severe as in adults, except the recurrent colicky abdominal pain seen in 40–80% of children [55,58,107].

It is important to note that attacks of laryngeal oedema can occur at any age and may be life-threatening [79]. For this reason, particularly where there is a family history, children should be tested at an early age. There are few data confirming the reference range for C1 INH in the very young. We would advocate testing both C4 and C1 INH to confirm the diagnosis in these circumstances.

**Long-term prophylaxis of attacks in children**

This is a relatively unexplored issue [58,111], and most references state that the use of antifibrinolytics and androgens are not recommended because of the serious side effects of these drugs [25,76].

Severe or life-threatening attacks of C1 INH deficiency are less common during childhood but they do occur. Although earlier reviews suggest prophylaxis is rarely required in children [15,55], this view is changing with increasing experience with tranexamic acid. Long-term prophylaxis is
justified not only in severely affected children, defined by attacks of laryngeal oedema and/or frequent (more than one every 3 months), recurrent attacks of abdominal pain causing distress and disability. In this situation, antifibrinolytics are preferred to androgens [54,58,111]. The individual minimal effective dose, irrespective of serum concentrations of C4 and/or C1 INH, for both antifibrinolytics and/or androgens used for long-term prophylaxis, has to be established and adjusted with growth. The use of prophylactic attenuated androgens in children is hardly ever indicated and if tranexamic acid is ineffective then regular infusions of C1 INH concentrate should be considered (Table 6).

**Attenuated androgens**

Attenuated androgens are associated with increased risk of androgenization, premature puberty, accelerated bone fusion with limited growth, liver disorders, atherogenesis and behavioural problems. The use of danazol in children [114,115], particularly its potential effect on development, and particularly in respect of emerging infections.

The appropriate therapeutic dose to be held will depend on the size of the child and should be agreed with the specialist.

**Antifibrinolytics in children**

Tranexamic acid at a dose of 50 mg/kg/day [54] or 1.5 g/day [52,55] has been used long-term with similar benefit and no side effects. Long-term administration of high dose EACA (12–24 g/day) in children was associated with side effects in all, but when the dose was adjusted for each child’s need (6 g/day and 12 g/day for <11-year-olds and >11-year-olds, respectively), the control of symptoms was still satisfactory without unpleasant side-effects [58].

It has been proposed that the long-term use of antifibrinolytics, by plasmin inhibition, could also predispose to arteriosclerosis [58,123]. This is of particular importance if long-term prophylaxis is to be started during childhood because several decades of treatment may be needed.

**C1 inhibitor concentrate in children**

C1 inhibitor concentrate has been used successfully for long-term replacement in selected adult patients [124], and more recently it has been shown to be superior to a placebo in a double-blind controlled study [76]. In an uncontrolled trial during long-term follow-up of 14 children with C1 INH deficiency, acute attacks in six children were treated with a single dose of 500 U of C1 INH concentrate (Immuno AG, Vienna, Austria) on 30 separate administrations. Progression of facial and laryngeal oedema was aborted 30–60 min after the infusion and disappeared gradually over the next 24–36 h. The dose had to be repeated after 60 min on only two occasions because laryngeal oedema continued to progress. Concentrations of C1 INH and C4, when measured 12 and 24 h after the infusion in two patients, showed an expected increase. None of the children required endotracheal intubation or tracheotomy, and no side effects were observed.

Based on the clinical benefit seen in these patients, a role for C1 inhibitor concentrate in long-term prophylaxis for children has been suggested [76], supporting the few earlier proposals [15,125]. In children home therapy is difficult because of technical problems with intravenous access. However, home availability of concentrate has been shown to reduce access time in adults [80] and home possession of concentrate should have similar benefits in children [76,125,126]. Advice should be given that at the earliest sign of an attack involving the upper airway, treatment and medical assistance should be sought. The disadvantages to this approach to the management are expense [117] and the possibility of viral transmission. Despite the lack of evidence of viral transmission with current pasteurized products caution is required when recommending any blood product, particularly in respect of emerging infections.

Abdominal oedema in children may be the major presenting symptom of an acute attack. The many other causes of abdominal pain in childhood need to be considered. The cardinal feature of abdominal oedema in these cases is severe abdominal pain, usually with vomiting, which lasts several hours and may mimic acute appendicitis. Early treatment of symptoms is effective and may reduce the requirement for further treatment. It may also lead to avoidance of inappropriate surgery. Therefore, home possession of C1 INH concentrate may be beneficial. It is important that a management care plan is in place for the patient, ensuring sufficient supply for use and for immediate replacement after use to ensure an adequate supply in case of further attacks. See 'Patient possession of C1 INH concentrate’ for further details on home possession of C1 INH.

The appropriate therapeutic dose to be held will depend on the size of the child and should be agreed with the specialist.

### Table 6. Major considerations regarding potential long-term prophylaxis for C1 INH deficiency in children

| Four attacks per year requiring possible admission and intravenous fluids should not be taken lightly |
| The use of attenuated androgens is hardly ever indicated as long-term prophylaxis prepuberty |

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We recommend monitoring schedules consisting of pretreat-
pies and the risk of failing to treat laryngeal angio-oedema.
of the product, the comparative risk with using other thera-
viral infection, given a clear explanation of the safety record
[130] (see Table 7).

The patient should be informed of the potential dangers of
danger, given a clear explanation of the safety record
of the product, the comparative risk with using other thera-
ies and the risk of failing to treat laryngeal angio-oedema.
We recommend monitoring schedules consisting of pretreat-
ment screening for hepatitis B, hepatitis C, alanine ami-
otransferase (ALT) and storage of serum and DNA. Six-
hepatitis C virus (HCV) was observed in these studies. None the less,
surveillance of patients treated with concentrate is essential

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The viral safety of C1 inhibitor concentrate

As with any blood product, viral safety is always a matter of
concern. There are reports of transmission of hepatitis C
virus (HCV) by non-virus inactivated C1 INH concentrates
used before 1985 [110,127,128]. Several studies confirmed
the safety of a heat treatment step in the production of a C1
inhibitor concentrate [25,76,128,129] and no transmission
of the human immunodeficiency virus, HCV or hepatitis G
virus (HGV) was observed in these studies. None the less,

The patient should be informed of the potential dangers of
danger, given a clear explanation of the safety record
of the product, the comparative risk with using other thera-
ies and the risk of failing to treat laryngeal angio-oedema.
We recommend monitoring schedules consisting of pretreat-
ment screening for hepatitis B, hepatitis C, alanine ami-
otransferase (ALT) and storage of serum and DNA. Six-
monthly liver function tests are recommended if concentrate
has been infused. Recombinant preparations of C1 INH, if
successful, would overcome many of these difficulties.

By analogy with the recommendations in haemophilic
patients [131], consideration should be given to vaccinate
patients who are not immune to hepatitis A or B and who
currently receive, or may require, blood products. Note that
the hepatitis A vaccine is not licensed for use in children
under the age of 1 year.

FFP is effective in the treatment of acute attacks [132,133]
and in short-term prophylaxis [101,134,135], but carries sig-
nificant risks of pathogen transmission, anaphylactoid rea-
tions, alloimmunization and excess intravascular volume
[15,25]. FFP is used when C1 INH is unavailable but is not
acceptable where emergency treatment is foreseeable or as
prophylactic treatment [109,136,137].

Service specification

Diagnosis

The laboratory diagnosis should be made only by a
CPA-approved laboratory with the input of a clinical
immunologist.

Table 7. Advice on use of C1 INH concentrate

C1 INH concentrate should only be given for severe attacks of swelling
where there is a risk of airway involvement, for severe attacks of
abdominal pain or uncontrolled disease
Liver function and viral status of these patients should be monitored
regularly and records kept of all infusions given
Patients should be fully informed of the potential risks and involved in
treatment decisions
Consideration should be given to vaccinate patients who are not
immune to hepatitis A or B

Treatment, local versus regional

Ensure that each region has nominated centres with an
immunologist and specialist nurse input. The centre(s) must
have a sufficient number of patients and expertise to com-
petently diagnose patients and competently develop individ-
ual management plans, written protocols and advice sheets
for patients. This should include advice on where and how to
seek assistance in the emergency situation, the appropriate
testing of relatives and general management of the condition.
Appropriate home therapy training should be available. Remote
patients will be monitored locally by the dermatol-
ogist or other designated physician following protocols with
reference back to the regional centre within the protocols.

Information

Patients should have written information on their condition,
its treatment, the side effects of treatment and a plan on how
to obtain emergency treatment.

Training infusion

- Immunology or equivalent specialist to manage home
therapy training programme.
- Immunology or equivalent specialist to liaise with accident
and emergency (A&E) departments and GPs regarding
treatment of acute attacks of C1 INH deficiency.
- Immunology or equivalent specialist to have a key role
in the ongoing education and support of the patient
with regard to all aspects of their HAE management
programme.
- Emergency care: A&E, primary care and home therapy.

A&E departments

A treatment plan or protocol for the management of patients
with C1 INH deficiency should be accessible in the depart-
ment. Long waiting-time in A&E is a major factor in disrup-
tion to work or education and quality of life, and deters
patients from seeking appropriate treatment. Protocols
should include mechanisms for prioritizing these patients,
for example nurse-led protocol-driven treatment with med-
ical review if necessary.

The immunology nurse/nurse specialist should liaise with
senior A&E medical and nursing staff to ensure staff have a
basic knowledge of C1 INH deficiency and are aware of
locally known patients with this diagnosis.

Senior medical and nursing staff should know how to
obtain and administer C1 INH concentrate – this should be
covered in the protocol.

A&E staff should be aware of how to access specialist
(immunology) team if advice is required – this should also
be covered in the protocol.
GPs

A treatment plan or protocol for the management of C1 INH deficiency should be sent to GP practices. GPs should be made aware of any C1 INH-deficient patient registered with their practice. If the patient intends to infuse concentrate at home, the patient’s GP should be informed. Appropriate emergency cover for any complications must be provided.

The immunology nurse/nurse specialist should liaise with the GP regarding concerns or problems with home therapy, or other relevant issues relating to management of C1 INH deficiency patients.

Outcome measures

In the context of clinical governance, the following are considered as suitable (i.e. measurable) topics for clinical audit.

- Number of acute attacks per patient per year.
- Quality of life scores.
- Attack/pain to needle time.
- Frequency of visits to A&E.
- Death.
- Side effects of treatment, such as abnormal liver function tests or liver ultrasound.
- Compliance of centres with guidelines.

Register of patients

In order to improve further the understanding of HAE and to improve service to patients it is recommended that regional units submit patients’ details to the European register. A form for this is provided on the internet at http://www.haeregister.org.

Patients’ perspective

The key aims of C1 esterase inhibitor-deficient patients

- For each C1 INH-deficient patient to be able to manage his/her symptoms proactively in such a way that they maintain personal safety and minimal disruption in living a healthy and productive life.
- The universal availability of effective C1 INH deficiency management for all patients.
- To avoid misdiagnosis, inappropriate treatment and unnecessary surgical procedures.

Achievement of key aims

This can be achieved by:

- the referral of all patients to a specialist who has experience of C1 INH deficiency treatment.
- Recognizing the provision and key role of the specialist nurse in educating and supporting C1 INH deficiency patients.
- Effective communication between the team involved in the individual patient’s care.
- Disseminating information to both health professionals and patients.
- Networking and information sharing between all the specialties treating C1 INH deficiency patients so that there is an agreed approach to the key issues of C1 INH deficiency management.

Outcomes

- The patient has an enhanced quality of life. He/she is more likely to maintain employment and contribute fully to the life of the community. This reduces the requirement for support.
- It has been well demonstrated in other chronic conditions that the informed patient who takes responsibility for their condition will make fewer demands on healthcare systems.
- The effectiveness of this approach would be shown in fewer visits to GPs, consultants, A&E departments, less use of ambulance services and less need for hospital in-patient treatment.

Acquired angio-oedema

The recent paper by Cicardi et al. [8] has shown importantly that in long-term follow-up (up to 24 years, median 8 years), the majority of cases was associated with lymphoproliferative disorders, predominantly monoclonal gammopathies of uncertain significance (MGUS). This is important because approximately 1% of MGUS per year progress to myeloma or a related disorder [138]. A minority of cases is associated with non-haematological malignancy, infection or autoimmune disorders. Where possible, treatment of the underlying pathology may lead to resolution of the disorder [30,32] (evidence level 3). Otherwise, the treatment is similar to HAE. Antifibrinolytics are more effective than attenuated androgens in this group [8] (evidence level 2). In the series of Cicardi et al. [8] therapy with C1 INH concentrate was necessary in 12 of 28 patients, of whom three became progressively resistant.

Disclaimer

While the advice and information in these guidelines is believed to be true and accurate at the time of going to press, the authors can accept no legal responsibility or liability for any omissions or errors that may be made.

Consultation process

The standards were developed by the project Advisory Group (see Table 8 for membership) with the project consultants. Consultation on draft standards was undertaken through:
• A 1-day facilitated workshop with representatives of national stakeholder organizations.
• Individual meetings with professional groups, voluntary sector organizations and regional groups of providers.
• Request for written comments from all individuals and organizations invited to the above events.
• Availability of draft standards on Primary Immunodeficiency Association (http://www.pia.org.uk) and Primary Immunodeficiency Network websites (http://www.ukpin.org.uk/News/C1-inhibitor.doc) with request for comment.

Declaration

The final version has been read and approved by all members of the advisory group (Table 8). Members of the advisory group were selected to provide a range of experience and expertise in immunology service provision. The group as a whole advised on the project and the consensus in general. M.M.G., R.J.L. and L.M. comprised the writing team. In addition, each group member co-drafted individual sections.

References

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Table 8. Membership of the advisory group

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