Disclaimer

This document is a general guide to appropriate practice, to be followed subject to the clinician’s judgment and the patient’s preference in each individual case. The guidelines are designed to provide information to assist decision-making and were reviewed in 2010.

These guidelines will be reviewed again in 2013.
1. INTRODUCTION

The management of patients with haemophilia A and haemophilia B and inhibitors is complex and must be managed by a Haemophilia Treatment Centre (HTC). Major issues with treatment are firstly, the control of acute haemorrhage and secondly the eradication of the inhibitor (immune tolerance induction - ITI). A systematic review of management of inhibitors in haemophilia A and haemophilia B has been published [1, 2].

2. TOLERISATION ADVISORY COMMITTEE

Management of inhibitor patients is complex. The Tolerisation Advisory Committee (TAC) is a subcommittee of AHCDO providing consultation on all cases of inhibitor patients and tolerisation. In general, all cases of proposed ITI in Australia should be discussed within the setting of the TAC.

3. GENERAL COMMENTS

Approximately 25% of patients with severe Haemophilia A develop inhibitors after initial treatment with factor VIII concentrates [3]. Approximately half of these appear transient and disappear with continued therapy with factor VIII. Therefore the diagnosis of an inhibitor, irrespective of titre, requires active monitoring and treatment to determine whether the inhibitor progresses to a high titre inhibitor (>5BU) or is transient.

After initial diagnosis of a low titre inhibitor in a patient with Haemophilia A, routine doses of factor VIII may be as much as doubled and the clinical response and inhibitor titre monitored. Subsequent poor clinical response or rising titre levels indicate a non-transient inhibitor. The diagnosis of a high titre inhibitor which does not respond to factor VIII should be treated with an alternative product.

There is an incidence of inhibitors in those with mild Haemophilia A following treatment with factor VIII concentrates and the guidelines will also apply to those
patients. DDAVP is occasionally used in mild to moderate haemophilia but is often ineffective [3].

The incidence of inhibitors in those with severe Haemophilia B is much less common (3%) but requires more careful individual management in a HTC. The immune based problems of anaphylaxis and nephrotic syndrome can occur in the setting of ITI as an additional complication [1].

4. CONTROL OF HAEMORRHAGE

4.1. Products Available

In Australia the main products available are

- factor VIII, either recombinant or plasma-derived
- recombinant factor VIIa
- activated prothrombin complex concentrate (eg FEIBA)
- prothrombin complex concentrate (eg prothrombinex-HT)
- antifibrinolytics (eg tranexamic acid)

Note: Porcine factor VIII is no longer being manufactured. A recombinant porcine factor VIII may soon be available

4.1.1. Factor VIII / Factor IX

In the case of life threatening haemorrhage infusion of factor VIII or factor IX in large doses can be used to swamp the inhibitor. This therapy is mostly used in patients with low responding inhibitors (inhibitor titre is <5 BU/mL after infusion of factor VIII). In patients who are high responders, this treatment may be effective for short periods of time provided the inhibitor level is less than 5 BU/mL. Factor VIII and factor IX levels should be observed to assess and monitor response. Anamnesis can occur 3- 5 days after therapy and make factor VIII ineffective. Patients with factor IX inhibitors can experience allergic reactions upon re- exposure to factor IX and administration should only be given in the hospital setting with access to facilities to manage anaphylaxis.
4.1.2. Recombinant Factor VIIIa

This product has been widely used in Australia and has proved to be highly effective in the management of bleeding episodes in patients with inhibitors to FVIII or FIX. Evidence in the literature suggests that it is effective in 79-92% of such episodes [4]. In addition, there is evidence that it is effective in over 90% of cases of surgery [4].

Recombinant factor VIIIa is infused as a bolus. Continuous infusion of recombinant factor VIIIa may reduce the quantity and cost of treatment but evidence is conflicting. A recent study suggests continuous infusion of 50 µg/kg/hr is effective in surgery [3]. Antifibrinolytics may be administered concurrently but may increase the risk of thrombosis in some patients. The standard adult dose of recombinant factor VIIIa is 90 µg/kg. Doses may need to be repeated every 2–3 hours until bleeding settles. Recent evidence suggests that larger doses (e.g. 270 µg/kg) maybe as effective as repeated doses of 90 µg/kg [5]. In children the mean half life is substantially reduced to 1.32 hours and thus higher doses of up to 200-250 µg/kg may be required.

Recently the use of rVIIIa as secondary prophylaxis has been approved for funding by the NBA. The regime of 90µg/kg daily for up to 3 months may be trialled in patients who experience a high incidence of bleeding (defined as four bleeds per month or repeated serious bleeding episodes e.g. retroperitoneal bleeding).

4.1.3. Activated Prothrombin Complex Concentrates (APCCs) eg FEIBA

Activated prothrombin complex concentrate, such as FEIBA, is effective in the treatment of 90% of bleeding episodes, and has been effective in the management of bleeding during major surgery [4]. An effective dose is 60-100 units/kg twice per day. The maximum daily dose of FEIBA is 200 units/kg/day. Antifibrinolytic agents, such as tranexamic acid, should not be administered concurrently with FEIBA. It should be noted that FEIBA contains small amounts
of factor VIII and therefore may cause elevation of inhibitor titres in some patients.
In patients who are having frequent bleeds, a trial of FEIBA as prophylaxis may be considered. The suggested dose is 75-100 units/kg three times a week.

4.1.4. Antifibrinolytic therapy
Antifibrinolytic therapy is particularly effective for mucosal bleeding episodes. The recommended dose of tranexamic acid is 80-100 mg/kg/day, with a standard dose being 1g, *qid* given orally (recommended paediatric dose is 35 mg/kg/8hr). An intravenous preparation is available in Australia through the SAS. Intravenous epsilon aminocaproic acid (EACA) is no longer produced and not available.

4.1.5. Immunomodulation
A number of immunomodulation strategies, including plasmapheresis to reduce the inhibitor titre and allow treatment with FVIII, have been suggested for patients who with inhibitors to FVIII and FIX presenting with acute haemorrhage. Specific immunoadsorption using the Malmo protocol [5] is not available in Australia.

4.2. Treatment Regimens for control of haemorrhage

4.2.1. Low titre inhibitor (<5 BU/ml), low responder
The recommended dose of factor VIII is 50-100 IU/kg repeated every 8-12 hours. Doses of factor IX of 100 – 200 IU/kg 12 – 24 hourly may be used in patients with low titre factor IX inhibitors and no or limited allergic reactions to factor IX who present with bleeding.

4.2.2. Low titre inhibitor (<5 BU/ml) *AND* history of high responder
Infusions of factor VIII or factor IX will likely cause an anamnestic rise of the levels of factor VIII inhibitor within 3-5 days rendering further therapy with factor VIII or factor IX ineffective. Recombinant factor VIIa or FEIBA should be considered to manage bleeding episodes in these patients.
4.2.3. High titre inhibitor (>5 BU/ml)
Recombinant factor VIIa or FEIBA should be considered to manage bleeding episodes in these patients.

4.2.4. Elective Major Surgery
Major surgery in patients with inhibitors carries a high degree of risk and should only be carried out in recognised HTCs after careful consultation and agreement with at least one other Australian haemophilia specialist. The TAC can be utilised for this purpose. All such discussions should be documented. It is recommended that a pharmacokinetic study be undertaken in low titre patients or patients undergoing tolerisation before surgery. The dosage regimen is based on the regimen for major bleeds.

Dental surgery and the insertion of IV access devices require 3-5 days of therapy and antifibrinolytics.

4.2.5. Emergency Major Surgery
The dosage regimen is based on the regimen for major bleeds. If time allows, there should be consultation and agreement with one other haemophilia specialist as in Elective Major Surgery. Any such discussion should be documented. The patient should be transferred to a recognised HTC as soon as practicable.

4.2.6. Home Therapy
AHCDO recommends that home therapy for patients with inhibitors is closely monitored with regular medical supervision.

5. TOLERISATION

5.1. Haemophilia A
Tolerisation should be considered in all patients with inhibitors to factor VIII. Immune tolerance induction (ITI) is defined as eradication of an inhibitor by high dose antigen exposure with or without immune modulation therapy. Tolerisation is an active procedure that may involve development of antibodies against IgG
factor VIII or factor IX antibodies. Intensive replacement therapy for immune
tolerance usually requires central venous access.

Despite decades of experience with ITI, little information from scientific studies is
available to guide clinicians in the management of ITI. The international ITI
study which is a randomised trial between high dose therapy (200 units/kg/day)
and low dose therapy (50 units/kg three times per week) was initiated in 2002
and stopped in November 2009 because of safety concerns and “futility”. It was
recognised that nearly 400 evaluable patients would be required to answer
questions regarding dose and it was deemed this would not achievable. There
were increased total haemorrhages, joint bleeds and muscle bleeds in the low
therapy arm compared to the high dose regime. No advice on dose can be given
from this trial.

Findings from an International Workshop on Immune Tolerance Induction have
recently been published[3].

A number of international and national registries of tolerisation offer information
on prognostic information for patients having ITI. In a meta-analysis of these
registries, two predictors of success for ITI were identified; historical peak
inhibitor titre and titre of inhibitor at initiation of ITI [3]. Based on this information
and from the report from the International Immune Tolerance study [6], the
following are noted

- Waiting for the inhibitor titre to fall to < 10 BU is possible prior to
  commencement of ITI. The median delay between the development of an
  inhibitor and the drop off in titre is 5 months. (reference).

- In a good risk patient there is no information to guide clinicians in dose
  (high or low dose).

- There is no information to support the choice of particular factor VIII
  products in good risk patients and in general, the patient should be
tolerised with the same product used at the time of developing an
inhibitor.
• In poor risk patients (historical titre > 200 BU, > 5 years from diagnosis, pre-ITI titre > 10 BU) evidence suggests higher dose regimes may be more successful (e.g. 200 IU/kg/day). Clinicians may consider the use of plasma derived factor VIII in these patients.

• Patients who show poor response to ITI (defined as showing a < 20% reduction in inhibitor titre for any 6 month period after the first 3 months of treatment) should be considered for second therapy. This includes changing to a FVIII product containing vWF, increasing dose or immune modulation such as anti-CD20 therapy.

• Successful ITI is defined as
  1. Negative inhibitor screen
  2. FVIII recovery of > 66% of predicted
  3. FVIII half life of > 6 hours after a 72 hour wash out period

5.2. Haemophilia B
ITI for patients with haemophilia B and inhibitors to FIX is difficult. Successful ITI in these patients is significantly less than in patients with haemophilia A. Additional allergic complications of anaphylaxis and development of nephritic syndrome in the setting of ITI is well described but poorly understood. Patients with large gene deletions of the FIX gene may be more at risk of developing this complication. There is limited information available to guide clinicians in the management of these patients. A number of scattered case reports describe the use of various forms of immune modulation to maximise the success of ITI in patients with haemophilia B.

No recommendations regarding ITI can be provided for patients with haemophilia B and inhibitors to factor IX. Patients who have ITI attempted should have regular assessments to exclude proteinuria and development of nephrotic syndrome [3].
REFERENCES

1. Di Michele DM. Inhibitor development in haemophilia B; an orphan disease in need of attention. BJH 2007;138:305-315.


