### 7.0 APPROPRIATE USE OF MANUFACTURED BLOOD PRODUCTS

<table>
<thead>
<tr>
<th>Policy</th>
<th>The Transfusion Medicine Service follows established guidelines for use of all blood products.</th>
</tr>
</thead>
</table>
| Reason | • Assist in the efficacious use of blood products  
• Improve patient safety by providing the appropriate product at the right dosing schedule | |
| Patient population | • As indicated in subsequent sections concerning individual products |
| Responsibilities of the Medical Director, Transfusion Medicine | • Be familiar with the appropriate use of these products  
• Assist in utilization management of these blood products by:  
  » Developing policies, processes and procedures to screen requests for these products to ensure that the most appropriate product is used in the right dose  
  » Promote education of clinical and other staff in the appropriate use of these products  
  » Assist clinicians when orders deviate from established guidelines and dose recommendations |
| Responsibilities of the treating physician | • Obtain any necessary authorization or special product release |
| Responsibilities of Transfusion Medicine Service staff | • Follow associated technical procedures as written  
• Consult with the Medical Director, Transfusion Medicine (or delegate) as indicated in procedures, as necessary based on technologist’s skills and experience, or on the basis of available clinical or laboratory information  
• Report all instances where these products were not given to a patient who met criteria  
• Provide any required request forms and/or contact information needed to obtain appropriate authorizations |
| Associated documents | Ontario guidelines have been developed for the following:  
• IVIG  
• Albumin  
Refer to: [www.transfusionontario.org](http://www.transfusionontario.org)  
National guidelines have been developed for the following:  
• Prothrombin Complex concentrates (PCC)  
• Solvent detergent treated plasma  
• Recombinant FVIIa  
Refer to: [www.nacblood.ca](http://www.nacblood.ca) |

### 7.1 APPROPRIATE USE OF PLASMA FRACTIONATED PRODUCTS

Plasma fractionated products include:  
• Albumin  
• Intravenous immunoglobulin (IVIG)  
• Subcutaneous immunoglobulin  
• Specific immune globulins (RHIG; HepBIG; VZIG)  
• Human derived clotting factor concentrates.( FEIBA ,PCC, FXI)  
• Recombinant clotting factors (FVII, VIII, IX)

### 7.2 INDICATIONS FOR THE USE OF ALBUMIN

Refer to the Ontario Albumin Administration Recommendations  
[http://www.transfusionontario.org/media/albumin%20guidelines_final_20120821.pdf](http://www.transfusionontario.org/media/albumin%20guidelines_final_20120821.pdf)
7.3 INDICATIONS FOR THE USE OF INTRAVENOUS IMMUNOGLOBULIN (IVIG)

In April 2012 the Ontario Ministry of Health and Long Term Care launched their IVIG strategy. For a toolkit and the associated guidelines, refer to [http://www.transfusionontario.org/media/IVIG%20Toolkit_COM_2012.pdf](http://www.transfusionontario.org/media/IVIG%20Toolkit_COM_2012.pdf)

General Pre-requisites and indications for IVIG Use:
1. A diagnosis must be confirmed for all orders.
2. For immune deficiency conditions, serum IgG levels must be clinically assessed to ensure optimum dosing.
3. For all other indications, IVIG should only be used when other, less expensive, equally safe and efficacious alternatives have failed.
4. There must be regular clinical outcome assessment.
5. For all proposed treatments or course of treatments with IVIG the MOHLTC IVIG Request Form (see below) shall be completed by the requesting physician.
6. All request forms must be reviewed for appropriate indication and dosage interval.
7. Detailed information on all aspects of IVIG Utilization Management can be found in the Intravenous Immune Globulin Toolkit prepared by ORBCoN.

Special Requests for Use in Conditions not on the list of Approved Medical Conditions for IVIG Use:
- Subject to screening at the hospital level:
  » IVIG user hospitals shall select the appropriate physician/committee to review, and where appropriate, approve requests for indications not listed on the MOHLTC IVIG Request Form
  » The physician appointed to serve as the approving physician (or delegate) shall sign the request form
  » On the request form under the heading “Other” the non-licensed indication shall be entered
- In the event of urgent treatment in a life-threatening situation, the request for IVIG shall be met immediately following verification of appropriate dose

Table 7.1 Approved Indications for IVIG Treatment
The clinical indication, dose and duration of therapy must be in accordance with the Ontario IVIG Utilization Management Guidelines. The list of recommended indications is as follows:

<table>
<thead>
<tr>
<th>Specialty</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Hematology</td>
<td>1.1. Fetal Neonatal Alloimmune Thrombocytopenia</td>
</tr>
<tr>
<td></td>
<td>1.2. Hemolytic Disease of the Fetus and Newborn</td>
</tr>
<tr>
<td></td>
<td>1.3. Idiopathic Thrombocytopenic Purpura Adult</td>
</tr>
<tr>
<td></td>
<td>1.4. Idiopathic Thrombocytopenic Purpura Pediatric</td>
</tr>
<tr>
<td></td>
<td>1.5. Post Transfusion Purpura</td>
</tr>
<tr>
<td>2. Neurology</td>
<td>2.1. Chronic Inflammatory Demyelinating Polyneuropathy</td>
</tr>
<tr>
<td></td>
<td>2.2. Guillain-Barre Syndrome</td>
</tr>
<tr>
<td></td>
<td>2.3. Multifocal Motor Neuropathy</td>
</tr>
<tr>
<td></td>
<td>2.4. Myasthenia Gravis</td>
</tr>
<tr>
<td>3. Dermatology</td>
<td>3.1. Dermatomyositis</td>
</tr>
<tr>
<td></td>
<td>3.2. Pemphigus Vulgaris and Variants</td>
</tr>
<tr>
<td>4. Rheumatology</td>
<td>4.1. Juvenile Dermatomyositis</td>
</tr>
<tr>
<td></td>
<td>4.2. Kawasaki Disease</td>
</tr>
<tr>
<td>5. Infectious Diseases</td>
<td>5.1. Staphylococcal toxic shock</td>
</tr>
<tr>
<td></td>
<td>5.2. Invasive Group A streptococcal fasciitis with associated toxic shock</td>
</tr>
<tr>
<td>6. Immunology</td>
<td>6.1. Primary Immune Deficiency</td>
</tr>
<tr>
<td></td>
<td>6.2. Secondary Immune Deficiency</td>
</tr>
<tr>
<td></td>
<td>6.3. High risk allogeneic stem cell transplantation</td>
</tr>
<tr>
<td>7. Solid Organ Transplant</td>
<td>7.1. Acute antibody mediated rejection</td>
</tr>
<tr>
<td></td>
<td>7.2. Kidney transplant from living donor</td>
</tr>
</tbody>
</table>

[http://www.transfusionontario.org/media/IVIG_Request_Form_R_31032012.pdf](http://www.transfusionontario.org/media/IVIG_Request_Form_R_31032012.pdf)
Indications for which IVIG is NOT recommended nor indicated, or is ineffective:

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Efficacy/Comment</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rheumatoid Arthritis</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
<tr>
<td>Inclusion Body Myositis</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
<tr>
<td>Chronic Fatigue Syndrome</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
<tr>
<td>Recurrent Spontaneous Abortion</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
<tr>
<td>In Vitro Fertilization/Implantation Procedures</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
<tr>
<td>Sepsis In Critical Care Patients</td>
<td>No large randomized controlled trials to confirm benefit</td>
<td>Not recommended</td>
</tr>
<tr>
<td>Autologous Bone Marrow/Stem Cell Transplant</td>
<td>No benefit</td>
<td>Not indicated</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
<tr>
<td>Amyotrophic Lateral Sclerosis</td>
<td>Ineffective</td>
<td>Not indicated</td>
</tr>
</tbody>
</table>

**IVIG dose calculation:**
- An IVIG Dose Calculator based on adjusted body weight is available to determine the appropriate dose for each individual patient.
- Available at [www.transfusionontario.ca](http://www.transfusionontario.ca) and as an application for installation on hand-held electronic devices.
- Based on patient height and weight.
- For IVIG for immunoglobulin replacement, use dose calculator for 1st dose and determine subsequent doses based on the IgG trough level.
- Preparations of immunoglobulin are available from Canadian Blood Services. The clinical indications for use are included in Table 7. Dosage is individualized, consult package insert.

**REFERENCES**
35. “Clinical Guidelines for Immunoglobulin Use.”
70. “Intravenous Immune Globulin.”
120. Shehata, N. 2010.
<table>
<thead>
<tr>
<th>Product</th>
<th>Clinical Indication</th>
<th>Standard Dose (Always refer to the product insert for most current information)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antihemophilic factor/ Von willebrand Factor complex Humate P® Wilate®</td>
<td>VonWillebrand’s disease when unresponsive to DDAVP</td>
<td>Minor bleed: either product 20–40 IU/kg q 12–24 hours Major Bleed: Humate P® 40–80 IU/kg q 12–24 hours Wilate® 40–60 IU/kg q 8–12 hours</td>
</tr>
<tr>
<td>Factor VIII concentrate*</td>
<td>Hemophilia A</td>
<td>• 1 U/kg produces 2% increase in factor VIII level • Half life 8–12 hours</td>
</tr>
<tr>
<td>Factor IX concentrate*</td>
<td>Hemophilia B (Christmas disease)</td>
<td>• 1 U/kg produces 1% increase in factor IX level • Half life 18–24 hours</td>
</tr>
<tr>
<td>Factor XIII Concentrate*</td>
<td>Congenital factor XIII deficiency</td>
<td>Consult package inserts</td>
</tr>
<tr>
<td>Antithrombin III</td>
<td>Antithrombin deficiency • Congenital deficiency • Heparin resistance in association with cardiovascular surgery</td>
<td>Refer to package insert.</td>
</tr>
<tr>
<td>Resource Manual for Medical Directors of Transfusion Medicine</td>
<td>Version 1, March 2013 <a href="http://www.transfusionontario.org">www.transfusionontario.org</a></td>
<td></td>
</tr>
</tbody>
</table>

### 7.0 APPROPRIATE USE OF MANUFACTURED BLOOD PRODUCTS

#### Rh Immune Globulin (RhIG)

- **Note:** Following any event at 20 weeks gestation or thereafter, feto-maternal testing should be performed to determine if additional doses of Rhig are required.

- **For prevention of RhD alloimmunization in at-risk RhD negative females**
  - **Pregnancy**
    - At 28 weeks gestation and post-partum with RhD positive infant
  - **Obstetrical**
    - Abortion -- therapeutic, spontaneous or threatened
  - Amniocentesis or chorionic villus sampling (CVS) <34 weeks gestation
  - Amniocentesis, CVS or other manipulation >34 weeks gestation
  - Additional sensitizing events (e.g. trauma, fall)
  - Post-transfusion of Rh D Positive red blood cells or platelets

- **Dose at 28 weeks, 1,500 IU or 300ug**
  - Dose post-partum 1,500 IU or 300ug, (may require additional doses as calculated following quantitation of feto-maternal hemorrhage)
  - 1,500 IU or 300ug

- **1,500 IU or 300ug**

- **600 IU or 120ug**

- **<12 weeks : 600 IU or 120ug**
  - ≥ 12 weeks: 1500 IU or 300ug

- **1,500 IU or 300ug for each 15mL red blood cells or 30 mL whole blood**

#### Varicella-Zoster Immune Globulin (VZIG)

- **Passive immunization to chickenpox in high risk exposed patients**

- **Dose at 28 weeks, 1,500 IU or 300ug**
  - 125 U/10kg to maximum of 625U, within 96 hours of exposure

#### Hepatitis B Immune Globulin (HBIG)

- **Passive immunization of exposed patients**

- **Dose at 28 weeks, 1,500 IU or 300ug**
  - Dose 0.06ml/kg immediately
  - Repeat in 1 month if not vaccinated

#### Tetanus Immune Globulin (TIG)

- **Passive immunization of exposed patients**

- **For dose, refer to package insert**

#### Prothrombin Complex Concentrates (Octaplex® and Beriplex®)

- **Treatment of major bleeding or in anticipation of urgent surgery in acquired deficiency of the prothrombin complex coagulation factors due to vitamin K antagonists or deficiency**

- **Effective half-life is only about 6 hours**
  - INR <3 – 1,000 IU
  - INR 3-5 – 2,000 IU
  - INR >5 (adults only) 3,000IU
  - Adjust for patients with extremes of body weight (<50kg, >90 kg)

- **C1 esterase inhibitor**

- **Treatment of hereditary angioedema in C1 esterase deficiency**

- **Refer to product insert**

#### Fibrinogen concentrate (riaSTAP™)

- **Congenital hypofibrinogenemia**

- **Refer to product insert**
### Table 7.3: Use of Products that Require approval through Health Canada Special Access Program or SAP

<table>
<thead>
<tr>
<th>Plasma Fractionated Product</th>
<th>Clinical Indication</th>
<th>Dose Information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factor VII concentrate</td>
<td>Congenital FVII deficiency</td>
<td>Refer to product insert</td>
</tr>
<tr>
<td>Factor XI concentrate</td>
<td>Congenital FXI deficiency</td>
<td>Refer to product insert</td>
</tr>
<tr>
<td>Factor XIII concentrate</td>
<td>Congenital FXIII deficiency</td>
<td>Refer to product insert</td>
</tr>
<tr>
<td>Protein C concentrate</td>
<td>Congenital or acquired deficiency of Protein C</td>
<td>Refer to product insert</td>
</tr>
</tbody>
</table>

#### REFERENCES


### 7.4 USE OF RECOMBINANT FACTOR VIIA, ERYTHROPOIETIN

#### Policy

The Transfusion Medicine Service follows established guidelines for use and dosage of recombinant products for the purposes outlined below.

The Medical Director, Transfusion Medicine has established a process to screen requests for recombinant products. This process includes creation and maintenance of a record of the patient response to therapy and outcome. All first time requests for recombinant factors must be approved by the Medical Director, Transfusion Medicine to address the issue of “off-label” use of recombinant factor VIIa.

#### Applies to

- Factor VIIa:
  - Control of bleeding in congenital factor VII deficiency
  - Patients with hemophilia A and B with coagulation factor inhibitors
  - Patients with acquired coagulation factor inhibitors refractory to medical therapy
  - Not recommended for treatment of bleeding in patients without the disorders listed above

#### Responsibilities of the Medical Director, Transfusion Medicine

- Be familiar with the availability and use of recombinant products and be aware that:
  - NAC guidelines recommend against off label use
  - Random controlled trials do not support off label use
  - Risk of adverse event is doubled over age 65 and tripled over age 75
- Understand the indications for and use of erythropoietin in the management of perioperative patients and for patients who refuse blood transfusion
- Ensure effective use of recombinant products by:
  - Screening requests for recombinant products
  - Promoting education of treating physicians and other health care professionals in the appropriate use of recombinant products
- Manage the inventory by:
  - Determining if the patient population served warrants holding a supply of these products as part of regular inventory, or should be requested from Canadian Blood Services on an ad hoc basis
  - Ensuring recirculation for expiring products in a timely fashion
Responsibilities of Transfusion Medicine Staff

- Follow associated technical procedures as written
- Respond promptly to requests where there life-threatening hemorrhage
- Insist that proper documentation is followed
- Order, receive and issue recombinant products
- Contact the Medical Director, Transfusion Medicine on receiving first-time requests for recombinant products
- Be aware that for all off-label requests for refractory bleeding:
  » NAC guidelines recommend against off label use
  » Random controlled trials do not support off label use
  » Risk of adverse event is doubled over age 65 and tripled over age 75

Conditions

- Recommended dosing for recombinant factor VIIa for:
  » Inhibitor patients - 90u/kg 2 hourly
  » Congenital factor VII deficiency – 15-30u/kg 2-8 hourly

Erythropoietin

Although erythropoietin is not distributed through the Transfusion Medicine Service, the Medical Director, Transfusion Medicine should be familiar with the drug and its indications for use in the peri-operative period and for patients who refuse blood transfusion.

REFERENCES