

<b>Section 7:</b>	<b>Biological Product Information</b>	<b>Standard #: 07.200</b>
<b>Created by:</b>	Province-wide Immunization Program Standards and Quality	
<b>Approved by:</b>	Province-wide Immunization Program Standards and Quality	
<b>Approval Date:</b>	January 1, 2017	<b>Revised:</b> February 4, 2019

<b>Botulism Antitoxin Heptavalent (Equine) Types A, B, C, D, E, F and G</b>	
<b>Manufacturer</b>	Emergent BioSolutions Canada Inc.
<b>Biological Classification</b>	Antitoxin
<b>Indications for Provincially Funded Vaccine</b>	<p><b>Treatment of botulism for individuals</b>– suspected or confirmed (administer immediately on suspicion of botulism - do not delay treatment waiting for lengthy clinical observations or confirmatory lab results).</p> <ul style="list-style-type: none"> <li>• Botulism Antitoxin is <b>not generally recommended for infants younger than one year of age</b>. Baby BIG® (Baby Botulism Immune Globulin (is the first-line therapy for naturally occurring infant botulism. See #07.201 Baby Botulism Immune Globulin. <ul style="list-style-type: none"> <li>○ Botulism Antitoxin Heptavalent may be considered for non-type A &amp; B infant botulism on a case-by-case basis.</li> </ul> </li> <li>• Eligibility will be determined though discussion amongst the treating physician, the zone Medical Officer of Health (MOH) and Alberta Health Chief Medical Officer of Health. <ul style="list-style-type: none"> <li>○ The zone MOH/MOH designate must notify the Office of the Chief Medical Officer of Health (OCMOH) by the fastest means possible of all cases in which botulism antitoxin is required.</li> <li>○ OCMOH is available on a 24 hour basis by pager at 780-638-3630.</li> <li>○ Once eligibility has been determined, OCMOH will authorize release of the product through either the Provincial Vaccine Depot or the Alberta Health Services (AHS) Calgary Vaccine Depot. It is not routinely stocked outside of these two sites.</li> </ul> </li> </ul> <p>For further information about the disease and reporting requirements refer to the <a href="#">AH Public Health Disease Management Guidelines – Botulism</a></p>
<b>Serology</b>	Blood (serum) should be collected to identify the specific toxin before antitoxin is administered; however the administration of the antitoxin should not be withheld pending the test results (refer to the <a href="#">AH Public Health Disease Management Guidelines – Botulism</a> )
<b>Schedule</b>	<p><b>Treatment:</b>  Infusion depends on age and weight. Refer to product monograph.  <a href="https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf">https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf</a></p>
<b>Preferred Use</b>	N/A
<b>Dose</b>	<p>Dose depends on age and weight. Refer to product monograph.  <a href="https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf">https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf</a></p> <p><b>Note:</b></p> <ul style="list-style-type: none"> <li>• This is a treatment product administered under the direction of a physician in an acute care setting.</li> <li>• Product is freezer stable and must be thawed prior to use. Refer to product monograph.</li> </ul>

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<b>Route</b>	This product is administered by slow intravenous infusion after dilution in normal saline based on dose recommendations in the product monograph. <a href="https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf">https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf</a>
<b>Contraindications/Precautions</b>	<p><b>Contraindications:</b></p> <ul style="list-style-type: none"> <li>• None as this is a vital indication due to life-threatening condition.</li> </ul> <p><b>Precautions:</b></p> <ul style="list-style-type: none"> <li>• Individuals who have received previous therapy with an equine-derived antivenom/antitoxin, or have known allergies to horses, or have asthma or get hay fever (seasonal allergies) may be at increased risk of hypersensitivity reactions and should only receive BAT if the benefits outweigh the risks.<sup>1</sup> Individuals should be closely monitored during and following administration.</li> <li>• Administer BAT in a setting with appropriate equipment, medication, including epinephrine, and personnel trained in the management of hypersensitivity, anaphylaxis, and shock.</li> <li>• Refer to Product Monograph <a href="https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf">https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf</a></li> </ul>
<b>Possible Reactions</b>	<p><b>Common:</b></p> <ul style="list-style-type: none"> <li>• Headache, nausea, pruritus and urticaria</li> <li>• Fever, chills, rash and edema</li> </ul> <p><b>Rare:</b></p> <ul style="list-style-type: none"> <li>• Allergic reaction</li> <li>• Infusion reactions (including chills, fever, headaches, nausea and vomiting). Monitoring required. Refer to product monograph.</li> <li>• Serum sickness (fever, urticarial or maculopapular rash, myalgia, arthralgia and lymphadenopathy may occur following botulism antitoxin administration typically 10-21 days after infusion.</li> <li>• Anaphylaxis</li> <li>• As with any immunization, unexpected or unusual side effects can occur. Refer to the product monograph for more detailed information of possible reactions and recommendations for management of those reactions.</li> </ul>
<b>Pregnancy</b>	<ul style="list-style-type: none"> <li>• There are no human or animal data to establish the presence or absence of risk associated with heptavalent BAT. Trivalent (A, B and E) BAT has been given to pregnant women without causing harm to mother or fetus. The benefit to the mother and the fetus from receiving heptavalent BAT for botulism should be weighed against the risk of harm from the treatment; decisions should be made on a case by case basis.</li> </ul>
<b>Lactation</b>	<ul style="list-style-type: none"> <li>• It is not known whether botulism antitoxin is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when botulism antitoxin is administered to a nursing mother. Botulism antitoxin heptavalent may be considered on a case by case basis.</li> </ul>
<b>Composition</b>	<p><b>Each single use vial (regardless of size or fill volume) contains a minimum antitoxin potency of:</b></p> <ul style="list-style-type: none"> <li>• Sterile solution of purified F(ab')<sub>2</sub> plus F(ab')<sub>2</sub>-related immune globulin fragments derived from equine plasma, containing antitoxin activity to botulinum neurotoxins A, B, C, D, E, F, G</li> <li>• 4500 U serotype A antitoxin</li> <li>• 3300 U serotype B antitoxin</li> <li>• 3000 U serotype C antitoxin</li> <li>• 600 U serotype D antitoxin</li> <li>• 0 5100 U serotype E antitoxin</li> <li>• 3000 U serotype F antitoxin</li> </ul>

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	<ul style="list-style-type: none"> <li>• 600 U serotype G antitoxin</li> <li>• No preservative</li> <li>• Clinically relevant non-medicinal ingredients               <ul style="list-style-type: none"> <li>○ 10% maltose</li> <li>○ 0.03% polysorbate 80</li> </ul> </li> </ul>
<b>Blood/Blood Products</b>	Equine horse serum
<b>Bovine/Porcine Products</b>	None listed in ingredient list
<b>Latex</b>	Does not contain latex
<b>Interchangeability</b>	N/A
<b>Administration with Other Products</b>	Must not be mixed with other medicinal products in a single container.
<b>Appearance</b>	Thawed product is a clear or slightly opalescent, liquid free of turbidity and foreign particles. Visually inspect the product for particulate matter and discoloration prior to administration. Do not use the solution if it is turbid or contains particles, other than a few translucent to white proteinaceous particulates.
<b>Storage</b>	<p>Storage:</p> <ul style="list-style-type: none"> <li>• Product is freezer-stable and must be stored at -15°C or colder. Do not refreeze.</li> </ul> <p>Administration:</p> <ul style="list-style-type: none"> <li>• Product must be thawed prior to use. Bring vial to room temperature prior to use. Refer to product monograph for product preparation instructions.  <a href="https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf">https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf</a></li> </ul> <p>Acceptable Temperature Management for BAT:</p> <ul style="list-style-type: none"> <li>• Once product has been thawed store in refrigerator between +2°C to +8°C.</li> <li>• Once the vial is opened, the preparation should be used immediately.</li> <li>• Do not re-freeze.</li> <li>• Do not microwave.</li> <li>• The product remains viable when stored at +2°C to +8°C as demonstrated by stability studies for a maximum of 38 months or until 48 months from the date of manufacture, whichever comes first.<sup>5</sup></li> </ul> <p><b>Note:</b> This does not imply or provide permission for the product to be <b>routinely</b> stored at +2°C to +8°C.<sup>5</sup></p> <ul style="list-style-type: none"> <li>• Date of manufacture: The date of manufacture is provided in the 'Certificate of Analysis' release letter which will be included with the product when it is shipped.</li> <li>• Do not use beyond the labeled expiry date.</li> <li>• Store in original packaging when possible to protect from light.</li> </ul>
<b>Vaccine Code</b>	BA
<b>Antigen Code</b>	BA
<b>Licensed for</b>	Botulism antitoxin is approved for sale in Canada and is currently only available via CMOH approval and release.
<p><b>Notes:</b> Botulism antitoxin is made from equine plasma, it may carry the risk of transmitting infectious agents, e.g., viruses. The equine plasma pools are screened for the presence of certain infectious agents and the manufacturing process for botulism antitoxin includes measures to inactivate and remove certain viruses. Despite these measures these products can potentially transmit disease. No cases of transmission of viral diseases have been associated with the use of botulism antitoxin. Refer to product monograph.</p>	

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### Related Resources:

- September 2018, special access program forms no longer required.
- **AHS-Imm-07.200-R01 (January 1, 2017)** Botulism Antitoxin Information Sheet

### References:

1. Alberta Health, Public Health and Compliance Division, Alberta Immunization Policy (2018, November 27). Botulism Antitoxin Heptavalent (Equine) Types A, B, C, D, E, F and G.
2. Alberta Health. Public Health Disease Management Guidelines - Botulism. In <https://open.alberta.ca/publications/botulism>
3. American Academy of Pediatrics. (2018-2021). Red book: 2018-2021 Report of the Committee on Infectious Diseases (31st ed.) Elk Grove, IL: Author.
4. Emergent BioSolutions Canada Inc. (2017 May 9) Botulism Antitoxin Heptavalent (A, B, C, D, E, F, G) – (Equine). Product Information. <https://emergentbiosolutions.com/sites/default/files/inline-files/BAT%20Product%20Monograph%20-%20English.pdf>
5. Emergent BioSolutions Canada Inc. (2018, October 31). Memo - Acceptable Temperature deviation for Botulism Antitoxin (BAT).