



## Bird's Eye Medical COVID-19 Monoclonal Antibody (mAb) EUA Treatment Referral: EVUSHELD

Patient Information	
<b>Name:</b> _____ <b>Sex:</b> M / F <b>Date of birth:</b> _____	
<b>Weight/BMI if high risk factor:</b> _____ <b>Phone:</b> _____ <b>Alt phone:</b> _____	
<b>Home Address:</b> _____	
<p><b>NOTE: For patients with moderate to severely compromised immune systems and not currently infected with COVID-19.</b> This product is available for use by Emergency Use Authorization (EUA) as pre-exposure prevention of COVID-19 for up to six months in people 12 years of age and older weighing at least 40 kg who are not currently infected with COVID-19 and who have had no known recent exposure to COVID-19 <b>and</b> have moderate to severely compromised immune systems due to a medical condition or immunosuppressive medications or treatments <b>and</b> may not mount an adequate immune response to COVID-19 vaccination <b>or</b> a history of severe adverse reactions to a COVID-19 vaccine and/or component(s) of those vaccines. This product is NOT a substitute for vaccination, nor authorized for the treatment of COVID-19 or for post-exposure prevention of COVID-19.</p> <p><b>NOTE on EVUSHELD dosing:</b> The FDA updated dosing guidelines of EVUSHELD from 150 mg each of tixagevimab and cilgavimab to 300 mg of each medication on 2/24/22. Per the updated guidance 6/2022, a complete dose of 300 mg each of tixagevimab and cilgavimab should be administered every 6 months. For those who received two separate doses of 150 mg of each medication, redosing should be 6 months after the last dose.</p>	
Treatment Criteria for Use (all fields must be completed to be eligible for treatment)	
<b>Treatment-qualifying high-risk condition(s):</b> <ul style="list-style-type: none"> <li><input type="checkbox"/> Active treatment for solid tumor and hematologic malignancies</li> <li><input type="checkbox"/> Receipt of solid-organ transplant and taking immunosuppressive therapy</li> <li><input type="checkbox"/> Receipt of chimeric antigen receptor (CAR)-T-cell or hematopoietic stem cell transplant (within 2 years of transplantation or taking immunosuppression therapy)</li> <li><input type="checkbox"/> Moderate or severe primary immunodeficiency (e.g., DiGeorge syndrome, Wiskott-Aldrich syndrome)</li> <li><input type="checkbox"/> Advanced or untreated HIV infection (people with HIV and CD4 cell count &lt;200mm<sup>3</sup>, history of an AIDS-defining illness without immune reconstitution, or clinical manifestations of symptomatic HIV)</li> <li><input type="checkbox"/> Active treatment with high-dose corticosteroids (i.e., ≥20mg prednisone or equivalent/day when administered for ≥2 weeks), alkylating agents, antimetabolites, transplant-related immunosuppressive drugs, cancer chemotherapeutic agents classified as severely immunosuppressive, tumor-necrosis (TNF) blockers, and other biologic agents that are immunosuppressive or immunomodulatory (e.g., B-cell depleting agents)</li> <li><input type="checkbox"/> Other medical conditions or factors: _____ (see CDC website for considerations)</li> </ul>	
Monoclonal Antibody Order:	
<input type="checkbox"/> <b>Repeat dosing; previous dose 6 months prior:</b> Tixagevimab 300 mg IM and Cilgavimab 300 mg IM <ul style="list-style-type: none"> <li><input type="radio"/> <b>Date of last dose:</b> _____ <i>please provide documentation of previous dose</i></li> </ul>	
<input type="checkbox"/> <b>First time dosing (all new EVUSHELD patients):</b> Tixagevimab 300mg IM and Cilgavimab 300 mg IM	
Prescriber Attestation	
<b>I affirm that my patient meets above criteria for use and has been:</b> <ol style="list-style-type: none"> <li>1. Informed that this mAb is an unapproved drug authorized for use under an <b>Emergency Use Authorization</b>.</li> <li>2. Informed of alternatives to receiving EVUSHELD.</li> <li>3. I have discussed this treatment option with the patient and patient is agreeable to this treatment.</li> </ol>	<b>I am aware that:</b> <ol style="list-style-type: none"> <li>1. EVUSHELD will be administered as a two separate, consecutive ventrogluteal IM injections. Patient will be monitored in the clinic for 1 hour post administration. Standard hypersensitivity reaction therapy will be provided as needed.</li> </ol>
Signature signifies agreement of the above orders / attestation.	
<b>Provider Name (print):</b> _____ <b>Office Number:</b> _____ <b>After hours phone:</b> _____	
<b>Provider Signature:</b> _____ <b>Date/Time:</b> _____ <b>NPI:</b> _____	
The prescribing health care provider and/or provider's designee should complete and submit a MedWatch form to FDA within 7 calendar days from the onset of a serious and unexpected adverse event that appears to be associated with the use of a monoclonal antibody.	

PLEASE FAX this form to 360.878.8330. If this is a repeat dose, send documentation of prior administration.

*If possible, attach copy of problem list, allergies, and medications.*

**Bird's Eye Medical will reach out to schedule your patient ASAP. For questions, please call 360-688-7044.**