Purpose: To meet the goal of administering FDA-Emergency Use Authorization casirivimab/imdevimab (REGEN-COV) to treat mild to moderate coronavirus disease, or for post-exposure prophylaxis (PEP) of COVID-19, in patients who are high risk for progression to severe COVID-19 and who meet the criteria set-forth by the Emergency Use Authorization Food and Drug Administration.

Policy: This standing order authorizes any North Carolina healthcare provider, in accordance with the conditions of their licensure and/or scope of practice to include subcutaneous injections, or pursuant to orders issued under North Carolina Executive Order 232, or as a covered person under the federal PREP Act functioning as monoclonal antibody providers to administer casirivimab/imdevimab (REGEN-COV) authorized by the FDA through an Emergency Use Authorization (EUA) and per conditions of this order.

### Casirivimab/Imdevimab (REGEN-COV) Administration

| Condition or Situation | Patients aged 12 years and older, weighing at least 40 kg (88.2 lb.), who present requesting and consent to treatment with monoclonal antibodies (casirivimab/imdevimab or REGEN-COV) for treatment of mild to moderate COVID-19 or for post-exposure prophylaxis to COVID-19, who self-attest to being at high risk for severe COVID-19 disease. Patients should have legal and decisional capacity to consent to treatment with monoclonal antibodies (casirivimab/imdevimab or REGEN-COV), in accordance with NC GS § 90-21.5. Casirivimab/imdevimab (REGEN-COV) can only be administered in settings in which healthcare providers have immediate access to medications to treat a severe infusion/injection or hypersensitivity reaction (such as anaphylaxis), and the ability to activate EMS, as necessary and according to local protocol. |

### Assessment Criteria

<table>
<thead>
<tr>
<th>Subjective</th>
<th>Treatment of Mild to Moderate COVID-19</th>
<th>Post-Exposure Prophylaxis of COVID-19</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Patient self-attests to positive results of SARS-CoV-2 viral testing <strong>AND</strong> 2. The patient presents within 10 days of symptom onset of COVID-19.</td>
<td>1. The patient self-attests that they are not <strong>fully vaccinated</strong> against COVID-19 or are not expected to <strong>mount an adequate immune response</strong> to complete COVID-19 vaccination <strong>AND</strong> 2. The patient self-attests that they are a <strong>close contact</strong> to an individual infected with COVID-19 or are at high risk of exposure to COVID-19 due to higher occurrence of infection in an institutional setting (for example, in nursing homes and correctional settings).</td>
<td><strong>Regarding repeat dosing:</strong> If the patient has an ongoing exposure to COVID-19 &gt;4...</td>
</tr>
</tbody>
</table>
In addition to meeting one of the above criteria, the patient self-attests to having a condition that would put them at high-risk for progression to severe COVID-19. Refer to the CDC’s review of People with Certain Medical Conditions for the most recent guidance on medical conditions that place a person at higher risk for severe illness with COVID-19. High risk conditions may include, but not be limited to:

1. Older age (for example, over the age of 65)
2. Obesity or being overweight (for example, BMI > 25; or if age 12-17, have BMI ≥ 85th percentile for their age and gender based on CDC growth charts)
3. Pregnancy
4. Chronic kidney disease
5. Diabetes
6. Immunosuppressive disease or immunosuppressive treatment
7. Cardiovascular disease (including congenital heart disease) or hypertension
8. Chronic lung diseases (for example, chronic obstructive pulmonary disease, moderate-to-severe asthma, interstitial lung disease, cystic fibrosis, and pulmonary hypertension)
9. Sickle cell disease
10. Neurodevelopmental disorders (for example, cerebral palsy) or other conditions that confer medical complexity (for example, genetic or metabolic syndromes and severe congenital anomalies)
11. Having a medical-related technological dependence (for example, tracheostomy, gastrostomy, or positive pressure ventilation unrelated to COVID-19)

Patients may self-attest they have another condition or factor that may put them at high risk for progression to severe COVID-19 disease that is not listed above.

If the patient presents with another condition or factor that is not listed above and the patient is uncertain if it may put them at high risk for progression to severe COVID-19 disease and therefore cannot self-attest to high risk, consult with the physician or
advanced practice provider (APP; nurse practitioner, certified nurse-midwife, or physician assistant) providing clinical supervision of the treatment facility/agency/service.

| Objective | 1. The patient is at least 12 years of age or older.  
|           | 2. The patient weighs at least 40 kg, or 88.2 lb. |

### Plan of Care

| Actions | 1. Review agency protocol for assessment and management of anaphylaxis before initiating treatment. Appropriate medical treatment and clinical staff able to manage immediate allergic reactions must be immediately available in the event an acute anaphylactic reaction occurs following administration reaction according to agency protocol.  
|         | 2. Prior to patients receiving casirivimab/imdevimab (REGEN-COV), provide and review the Fact Sheet for Patients, Parents and Caregivers EUA of REGEN-COV for COVID-19.  
|         | 3. Before administering casirivimab/imdevimab (REGEN-COV) or participating in any patient care activities, don appropriate personal protective equipment (PPE) per CDC guidelines to protect against the transmission of COVID-19. |

### Precautions: Patient Monitoring

The patient should be clinically monitored during and after administration of casirivimab/imdevimab (REGEN-COV). After administration is complete, the patient should be monitored for a minimum of 1 hour. During this time, the nurse, EMS personnel, or other individuals who are trained and supervised by clinical staff shall observe for signs and symptoms of a hypersensitivity reaction (anaphylaxis) or infusion/injection related reaction. These may include:

1. Fever  
2. Difficulty breathing  
3. Reduced oxygen saturation  
4. Chills  
5. Nausea  
6. Arrhythmia (such as atrial fibrillation, tachycardia, or bradycardia)  
7. Chest pain or discomfort  
8. Weakness  
9. Altered mental status  
10. Headache  
11. Bronchospasm  
12. Hypotension  
13. Hypertension  
14. Angioedema  
15. Throat irritation  
16. Rash (urticaria)  
17. Pruritus  
18. Myalgia  
19. Vasovagal reaction
Treatment

Subcutaneous Route-Initial Dose

1. Prepare 600 mg casirivimab and 600 mg imdevimab according to manufacturer instructions using aseptic technique into FOUR separate syringes. Casirivimab and imdevimab can be supplied as either a co-formulated package (REGEN-COV), or individually packaged. Refer to the Fact Sheet for Health Care Providers EUA of REGEN-COV for the most updated guidance on medication preparation for casirivimab/imdevimab.

Preparation of Casirivimab/Imdevimab for Initial Subcutaneous Injection

<table>
<thead>
<tr>
<th>Prepare 600 mg of Casirivimab and 600 mg of Imdevimab</th>
<th>Prepare 4 Syringes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Co-Formulated Vial</strong></td>
<td><strong>Withdraw 2.5 mL solution per syringe into FOUR separate syringes</strong></td>
</tr>
<tr>
<td><strong>Individual Vials</strong></td>
<td><strong>Casirivimab:</strong> Withdraw 2.5 mL solution per syringe into TWO separate syringes</td>
</tr>
<tr>
<td></td>
<td><strong>Imdevimab:</strong> Withdraw 2.5 mL solution per syringe into TWO separate syringes</td>
</tr>
</tbody>
</table>

2. Replace the 21-gauge transfer needle used for medication preparation with a 25-gauge or 27-gauge needle for subcutaneous injection. The recommended needle size for subcutaneous injections is 5/8”.

3. Administer the subcutaneous injections consecutively, each at a different injection site, into the thigh, back of the upper arm, and different quadrants of the abdomen, except for 2 inches (5 cm) around the navel. The waistline should be avoided.
   a. When choosing injection sites, avoid skin that is tender, damaged, bruised, or scarred.
   b. Follow local protocol for administering a subcutaneous injection.

If the patient is showing signs of anaphylaxis or an infusion/injection related reaction during or after administration; stop treatment, implement medical emergency protocols and immediately notify the physician or APP providing clinical supervision of the treatment facility/agency/service.
Subcutaneous Route - Repeat Dosing

1. Verify that the patient is presenting for repeat dosing of casirivimab/imdevimab (REGEN-COV) due to ongoing exposure to COVID-19 and that at least 4 weeks has passed since their last treatment.
2. Prepare 300 mg casirivimab and 300 mg imdevimab according to manufacturer instructions using aseptic technique into TWO separate syringes. Casirivimab and imdevimab can be supplied as either a co-formulated package (REGEN-COV), or individually packaged. Refer to the Fact Sheet for Health Care Providers EUA of REGEN-COV for the most updated guidance on medication preparation for casirivimab/imdevimab.

Preparation of Casirivimab/Imdevimab for Repeat Subcutaneous Injection

<table>
<thead>
<tr>
<th>Prepare 300 mg of Casirivimab and 300 mg of Imdevimab</th>
<th>Prepare 2 Syringes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co-Formulated Vial</td>
<td>Withdraw 2.5 mL solution per syringe into TWO separate syringes</td>
</tr>
<tr>
<td>Individual Vials</td>
<td>Casirivimab: Withdraw 2.5 mL solution per syringe into ONE separate syringe</td>
</tr>
<tr>
<td></td>
<td>Imdevimab: Withdraw 2.5 mL solution per syringe into ONE separate syringe</td>
</tr>
</tbody>
</table>

3. Replace the 21-gauge transfer needle used for medication preparation with a 25-gauge or 27-gauge needle for subcutaneous injection. The recommended needle size for subcutaneous injections is 5/8”.
4. Administer the subcutaneous injections consecutively, each at a different injection site, into the thigh, back of the upper arm, or different quadrants of the abdomen, except for 2 inches (5 cm) around the navel. The waistline should be avoided.
   a. When choosing injection sites, avoid skin that is tender, damaged, bruised, or scarred.
   b. Follow local protocol for administering a subcutaneous injection.

Follow-up

1. Provide the patient with COVID-19 Antibody Therapy Discharge Instructions and review it with them.
2. For patients who are receiving post-exposure prophylaxis for exposure to COVID-19:
   a. If the patient will have ongoing exposure to an individual with COVID-19, or is at high risk of exposure to an individual infected with COVID-19 due to high occurrence in an institutional setting (e.g., nursing homes or correctional facilities), advise the patient they should continue to
NC State Health Director’s Statewide Standing Order
for Subcutaneous Administration of Casirivimab/Imdevimab (REGEN-COV) Monoclonal Antibodies
September 13, 2021

<table>
<thead>
<tr>
<th>Contraindications for Use of this Order</th>
<th>Do not administer casirivimab/imdevimab (REGEN-COV) monoclonal antibody treatment to patients that:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Have previous severe hypersensitivity reaction, such as anaphylaxis, to casirivimab/imdevimab (REGEN-COV) or to any ingredient of casirivimab/imdevimab (REGEN-COV).</td>
</tr>
<tr>
<td>4.</td>
<td>Require an increase in baseline oxygen flow rate due to COVID-19 for patients on chronic oxygen therapy due to underlying non-COVID-19 related morbidity.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Criteria or Circumstances for Notifying the Physician or Advanced Practice Provider (APP)</th>
<th>Notify the physician/APP if:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>The patient desires treatment or post-exposure prophylaxis with casirivimab/imdevimab (REGEN-COV) but is uncertain if they meet the assessment criteria for use.</td>
</tr>
<tr>
<td>2.</td>
<td>The patient exhibits signs of a hypersensitivity reaction (anaphylaxis) or an infusion/injection-related reaction. In this instance, stop treatment; initiate emergency medical protocols and notify the physician/APP providing clinical supervision of the treatment facility/agency/service.</td>
</tr>
<tr>
<td>3.</td>
<td>Notify the physician/APP from the organization providing clinical supervision of the treatment facility/agency/service at any time there are questions or problems with carrying out this standing order.</td>
</tr>
</tbody>
</table>

Approved by: Elizabeth Cuervo Tilson, MD, MPH
NPI: 1760540421

Date approved: 9-13-21

This order is effective immediately upon signing and may be revised or revoked by the State Health Director according to his/her discretion. Legal Authority: Executive Order Number 232