AvMed

MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST*

Directions: The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-877-535-1391</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. If information provided is not complete, correct, or legible, authorization can be delayed.

For Medicare Members: Medicare Coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determination (NCD) and Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable. They can be found at: <u>https://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx</u>. Additional indications may be covered at the discretion of the health plan.

Drug Requested: Cinqair® IV (reslizumab) (J2786) (Medical)

MEMBER & PRESCRIBER INFORMATION: Authorization may be delayed if incomplete.

Member Name:	
Member AvMed #:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	
Office Contact Name:	
Phone Number:	
NPI #:	
DRUG INFORMATION: Authoriz	ation may be delayed if incomplete.
Drug Name/Form/Strength:	
Dosing Schedule:	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight (if applicable):	Date weight obtained:

□ Standard Review. In checking this box, the timeframe does not jeopardize the life or health of the member or the member's ability to regain maximum function and would not subject the member to severe pain.

Recommended Dosage: Dosage 3mg/kg <u>once every 4 weeks</u> by intravenous infusion over 20 - 50 minutes

*AvMed considers the use of concomitant therapy with Cinqair[®], Dupixent[®], Fasenra[®], Nucala[®], Tezspire[™] and Xolair[®] to be experimental and investigational. Safety and efficacy of these combinations have <u>NOT</u> been established and will <u>NOT</u> be permitted. In the event a member has an active Dupixent[®], Fasenra[®], Nucala[®], Tezspire[™] or Xolair[®] authorization on file, all subsequent requests for Cinqair[®] will <u>NOT</u> be approved.

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CLINICAL CRITERIA: Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied. (Trials will be verified using pharmacy claims and/or submitted chart notes.)

Initial Authorization: 12 months

- □ Prescribed by or in consultation with an allergist, immunologist or pulmonologist
- □ Member is 18 years of age or older
- □ Member has been diagnosed with severe eosinophilic phenotype defined by a baseline (pre-Cinqair[®]) peripheral blood eosinophil level of \geq 400 cells/microliter
- □ Member is currently being treated with <u>ONE</u> of the following unless there is a contraindication or intolerance to these medications and must be compliant on therapy <u>for at least 90 consecutive days</u> within a year of request:
 - □ High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) <u>AND</u> an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)
 - One maximally dosed combination ICS/LABA product (e.g., Advair[®] (fluticasone propionate/salmeterol), Dulera[®] (mometasone/formoterol), Symbicort[®] (budesonide/formoterol))
- □ Member has experienced <u>ONE</u> of the following (check box that applies):
 - More than 2 exacerbations requiring additional medical treatment (e.g., an increase in oral corticosteroid dose, emergency department, urgent care visits or hospitalizations) within the past 12 months
 - □ Any prior intubation for an asthma exacerbation
- □ Member has a baseline forced expiratory volume (FEV1) < 80% predicted normal submitted within year of request
- Provider must submit member blood eosinophil count collected after a trial and failure of at least 90 consecutive days of therapy with high dose inhaled corticosteroids <u>AND</u> long-acting inhaled beta-2 agonist. A failure of these medications is defined as a blood count > 400 cells/microliter (submit labs collected within the past 12 months)

Eosinophil count: _____

Date:

<u>Reauthorization</u>: 12 months. Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied. (Trials will be verified using pharmacy claims and/or submitted chart notes.)

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- □ Member has experienced a sustained positive clinical response to Cinqair[®] therapy as demonstrated by at least <u>ONE</u> of the following (check all that apply; chart notes must be submitted):
 - □ Increase in percent predicted Forced Expiratory Volume (FEV1) from baseline (pre-treatment)
 - **□** Reduction in the dose of inhaled corticosteroids required to control asthma
 - **□** Reduction in the use of oral corticosteroids to treat/prevent exacerbation
 - Reduction in asthma symptoms such as chest tightness, coughing, shortness of breath or nocturnal awakenings
- □ Member is currently being treated with <u>ONE</u> of the following unless there is a contraindication or intolerance to these medications:
 - □ High-dose inhaled corticosteroid (ICS) (e.g., greater than 500 mcg fluticasone propionate equivalent/day) <u>AND</u> an additional asthma controller medication (e.g., leukotriene receptor antagonist, long-acting beta-2 agonist (LABA), theophylline)
 - One maximally dosed combination ICS/LABA product (e.g., Advair[®] (fluticasone propionate/salmeterol), Dulera[®] (mometasone/formoterol), Symbicort[®] (budesonide/formoterol))

Medication being provided by (check applicable box(es) below):

Location/site of drug administration:

NPI or DEA # of administering location: _____

OR

D Specialty Pharmacy - PropriumRx

For urgent reviews: Practitioner should call AvMed Pre-Authorization Department if they believe a standard review would subject the member to adverse health consequences. AvMed's definition of urgent is a lack of treatment that could seriously jeopardize the life or health of the member or the member's ability to regain maximum function.

Use of samples to initiate therapy does not meet step edit/ preauthorization criteria. *<u>Previous therapies will be verified through pharmacy paid claims or submitted chart notes.</u>*