DESCRIPTION:

Prolastin-C®, Aralast NP®, Zemaira®, and Glassia® are all FDA approved agents for use as replacement therapy in congenital alpha-1 antitrypsin (AAT) deficiency with clinical emphysema. AAT deficiency is a hereditary recessive genetic disorder that increases the risk of chronic obstructive pulmonary disease (COPD), especially emphysema and chronic bronchitis. People with alpha-1 antitrypsin deficiency are at risk of degeneration of lung function, which may significantly affect quality of life and life expectancy.

AAT protects the delicate tissues of the lung by inhibiting the destructive action of an enzyme called neutrophil elastase. Neutrophil elastase is released by white blood cells, and its primary function is to digest bacteria and other foreign particles in the lungs. When circulating levels of AAT drop below a minimal protective level the alveolar walls are damaged from excess neutrophil elastase. When a person with AAT deficiency inhales irritants or contracts a lung infection, the neutrophil elastase released in the lungs continues to act uncontrolled, leading to destruction of healthy lung tissue. Patients with this disorder develop early onset panacinar emphysema (i.e. affecting all parts of the lobules). They are also at risk for developing chronic liver disease (hepatitis, cirrhosis), panniculitis (an inflammation of the layer of fat beneath the skin) and vasculitis. AAT deficiency represents about 3% of all emphysema cases reported in the United States.

POLICY:

Based upon our criteria and review of the peer-reviewed literature, treatment with Prolastin-C®, Aralast NP®, Zemaira®, and Glassia® administered in accordance with FDA guidelines, has been medically proven to be effective and therefore, appropriate if all of the following criteria are met:

1. Patient must be followed by and have a prescription written by a pulmonologist AND
2. Patient must currently be a non-smoker documented by a negative cotinine urine test.
   a. If using nicotine replacement products but no longer smoking, then urine anabasine measurements should also be ordered and must be negative AND
3. Patient must have one of the high-risk genotypes (such as PiZZ, PiSZ, PiZ(null), Pi(null,null), Pi(malton,malton), Pi(Siiyama,Siiyama)) or a dysfunctional AAT protein (such as PiF or Pi Pittsburg genotypes) as they are at the greatest risk for developing panacinar emphysema [please see exclusive criteria if patient has a PiMZ genotype] AND
4. Treatment should only be initiated when patient’s alpha 1-antitrypsin (AAT) levels are less than 11 micromol/L OR <57 mg/dl by nephelometry AND have documented evidence of emphysema as FEV1< 65% of predicted value.
   - Augmentation therapy is NOT recommended for patients without symptomatic emphysema
For those who meet the AAT level but have FEV1 > 65%, discussion with pulmonologist regarding potential benefits of therapy with consideration of cost (there is currently lack of evidence for benefit in this group. Factors such as age, rapid decline in FEV1, decreasing diffusing capacity, or progression of emphysema on imaging should be considered) AND

5. Patients should demonstrate 1 or more of the following: signs of significant lung disease such as chronic productive cough or unusual frequency of lower respiratory infection, airflow obstruction, accelerated decline of FEV1 or chest radiograph or CT scan evidence of emphysema, especially in the absence of a recognized risk factor (smoking, occupational dust exposure, etc.) AND

6. Patient must have had a trial and failure of Prolastin-C or have a contraindication to this therapy before treatment with Aralast NP, Zemaira or Glassia will be approved.

7. Approval dates are based on the patient’s contract. Please see chart in the policy guideline

8. Approved dosage for all of the drugs listed above is 60mg/kg IV infusion once weekly (Prolastin-C, Glassia, Aralast NP and Zemaira.). Doses higher than this are not recommended by clinical guidelines and therefore will not be covered.

POLICY GUIDELINES:

1. Unless otherwise stated above within the individual drug criteria, approval time periods are listed in the table below.

   a. Continued approval at time of recertification will require documentation that the drug is providing ongoing benefit to the patient in terms of improvement or stability in disease state or condition. Such documentation may include progress notes, imaging or laboratory findings, and other objective or subjective measures of benefit which support that continued use of the requested product is medically necessary. Also, ongoing use of the requested product must continue to reflect the current policy’s preferred formulary. Recertification reviews may result in the requirement to try more cost-effective treatment alternatives as they become available (i.e.; generics, biosimilars, or other guideline-supported treatment options). Requested dosing must continue to be consistent with FDA-approved or off-label/guideline-supported dosing recommendations.

Guidelines for approval time periods

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<tr>
<th>Line of Business</th>
<th>Medical Initial approval</th>
<th>Medical Recertification</th>
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<tr>
<td>Medicaid Managed Care (MMC) / Child Health Plus (CHP)</td>
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<td>12 months</td>
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<td>Outpatient Hospital – 6 months</td>
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<td>Outpatient Hospital – 6 months</td>
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2. This policy is applicable to drugs that are included on a specific drug formulary. If a drug referenced in this policy is non-formulary, please reference the Coverage Exception Evaluation Policy for All Lines of Business Formularies policy for review guidelines.

3. For contracts where Insurance Law § 4903(c-1), and Public Health Law § 4903(3-a) are applicable, if trial of preferred drug(s) is the only criterion that is not met for a given condition, and one of the following circumstances can be substantiated by the requesting provider, then trial of the preferred drug(s) will not be required.

   • The required prescription drug(s) is (are) contraindicated or will likely cause an adverse reaction or physical or mental harm to the member;
   • The required prescription drug is expected to be ineffective based on the known clinical history and conditions and concurrent drug regimen;
   • The required prescription drug(s) was (were) previously tried while under the current or a previous health plan, or another prescription drug or drugs in the same pharmacologic class or with the same mechanism of action was (were) previously tried and such prescription drug(s) was (were) discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event;
   • The required prescription drug(s) is (are) not in the patient’s best interest because it will likely cause a significant barrier to adherence to or compliance with the plan of care, will likely worsen a comorbid condition, or will likely decrease the ability to achieve or maintain reasonable functional ability in performing daily activities;
   • The individual is stable on the requested prescription drug. The medical profile of the individual (age, disease state, comorbidities), along with the rational for deeming stability as it relates to standard medical practice and evidence-based practice protocols for the disease state will be taken into consideration.
   • The above criteria are not applicable to requests for brand name medications that have an AB rated generic. We can require a trial of an AB-rated generic equivalent prior to providing coverage for the equivalent brand name prescription drug.

4. Patients MUST have clinically demonstrable panacinar emphysema

5. Patients with emphysema due to AAT deficiency should be maintained on regimens similar to those patients with emphysema not associated with AAT deficiency, including: maximum doses of beta-adrenergic bronchodilators, inhaled corticosteroids, anticholinergics and antibiotics, when appropriate. Patients should also have vaccinations against influenza and pneumococcus and supplemental oxygen therapy when indicated.

6. Treatment will only be covered when administered as an IV infusion.

7. Safety and effectiveness in children have not been established

EXCLUSIVE CRITERIA:
The use of alpha-1 Antitrypsin therapy will not be covered in any of the following situations:

1. Active smokers
2. Current non-smokers who start smoking after initial approval can be denied further treatment
3. Treatment of cystic fibrosis
4. Liver transplant recipients
5. Treatment of liver disease due to alpha-1 Antitrypsin deficiency
6. Augmentation therapy in general will not be granted for PiMZ heterozygotes or homozygous normal genotype (i.e. PiMM). Please note, commercial genotyping may identify a PiZ(null) heterozygote as PiMZ; because in the absence of an S or Z allele, most laboratories will assume

Proprietary Information of Health Plan
the second allele is M. If a patient with a documented PiMZ genotype and has an unusually low AAT level, this possibility should be considered.

CODES:
Eligibility for reimbursement is based upon the benefits set forth in the member’s subscriber contract. Codes may not be covered under all circumstances. Please read the policy and guidelines statements carefully.
Codes may not be all inclusive as the AMA and CMS code updates may occur more frequently than policy updates.

Code Key:
Experimental/Investigational = (E/I),
Not medically necessary/appropriate = (NMN).

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HCPCS:
J0256 Aralast NP, Prolastin-C, Zemaira 10mg per unit
J0257: Glassia 10mg per unit

Unit Threshold = 700

UPDATES:

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