What is Alpha-1 Antitrypsin Deficiency?

Alpha-1 antitrypsin deficiency is an inherited form of emphysema (em-fuh-ZEE-muh). People with the condition, also known as AAT Deficiency or alpha-one antitrypsin deficiency, do not have enough of a protein called alpha-1 antitrypsin (AAT) in their blood. This protein is made in the liver, and it protects the lungs so they can work normally.

Without enough AAT, the lungs can become damaged by chemical enzymes in the tissue that cause emphysema. Alpha-one antitrypsin deficiency also can also cause liver damage.

What is emphysema?

Emphysema is a condition that involves damage to the walls of the air sacs (alveoli) of the lung. Normally there are more than 300 million alveoli in the lung. These alveoli are stretchy and springy, like little balloons. Like a balloon, it takes effort to blow up normal alveoli; however, it takes no energy to empty the air sacs because they spring back to their original size.

In emphysema, the walls of some of the alveoli have been damaged. When this happens, the alveoli lose their stretchiness and trap air. Since it is difficult to push all of the air out of the lungs, the lungs do not empty easily and therefore contain more air than normal. This is called air trapping and causes hyperinflation in the lungs. In alpha-one antitrypsin deficiency, emphysema occurs more in the lower parts of the lungs than the upper parts.

The combination of constantly having extra air in the lungs and the extra effort needed to breathe causes a person to feel short of breath. Airway obstruction occurs in emphysema because the alveoli that normally help keep the airways open cannot do so during inhalation or exhalation. Without their support, the breathing tubes collapse, causing blockage (obstruction) to the flow of air.

What causes alpha-one antitrypsin deficiency?

Alpha-one antitrypsin deficiency is an inherited condition. Every person inherits two AAT genes—one from each parent. Inheriting two abnormal AAT genes causes alpha-one antitrypsin deficiency deficiency. A person who inherits only one abnormal gene is an alpha-one antitrypsin deficiency “carrier.” While a carrier’s AAT levels may be lower than normal, the risk of major health problems is much less than in a person with two abnormal genes.

Does everyone with two abnormal alpha-one antitrypsin deficiency genes develop disease?

Not everyone who inherits two abnormal AAT genes gets emphysema and/or liver disease. Some people never have symptoms. Some have only mild symptoms. Other people can have severe lung problems, liver problems, or both. Smoking is known to make lung disease worse if you have alpha-one antitrypsin deficiency.

What are the signs and symptoms of alpha-one antitrypsin deficiency?

Alpha-one antitrypsin deficiency lung symptoms are most often to appear after age 30, but may emerge earlier or much later. The first symptom is usually shortness of breath during daily activities. Other symptoms include wheezing and decreased ability to exercise.

How is alpha-one antitrypsin deficiency diagnosed?

Alpha-one antitrypsin deficiency can be detected through blood tests or a new test that painlessly takes a sample of DNA from the cells inside your mouth. One type of blood test measures the body’s level of AAT. If the AAT level is lower than normal, your health care provider may order a genotype or a phenotype blood test. These genetic analysis tests look at the amount and type of AAT being produced and compare it with normal patterns.

Alpha-one antitrypsin deficiency testing is
recommended for certain groups of people, including those who have:

- Family history of alpha-one antitrypsin deficiency
- Early-onset emphysema (less than 45 years old)
- Emphysema without an obvious risk factor such as smoking or occupational exposure to a substance known to cause the disease
- Emphysema that is worse at the bottom of the lungs
- Difficult to control asthma
- Recurrent pneumonia or bronchitis
- Unexplained liver disease
- Patients diagnosed with COPD should consult with their health care provider to see if they would benefit from being tested for alpha-one antitrypsin deficiency.

The health care provider also may recommend tests including a chest X-ray, a lung function test (a breathing test to find out how your lungs function compared with people with normal lungs), an arterial blood gas (measuring the level of oxygen in the blood) and liver function tests.

What are the treatments for alpha-one antitrypsin deficiency?

At this time, there is no cure for Alpha-one antitrypsin deficiency, but there are treatments that can improve symptoms. Your health care provider may prescribe medications such as bronchodilators or inhaled steroids to help open your airways. Your health care provider also may recommend pulmonary rehabilitation to improve your breathing. (For additional information on pulmonary rehabilitation, see ATS Fact Sheets listed in Resources) Patients with severe alpha-one antitrypsin deficiency may be candidates for a lung transplant. A treatment called AAT augmentation therapy, which may slow down or stop the destruction of lung tissue, may also be prescribed. This treatment increases the level of AAT in the blood. It is given intravenously (through a vein), and is usually given once a week for life.

Can I protect myself from lung damage if I have alpha-one antitrypsin deficiency?

If you have been diagnosed with alpha-one antitrypsin deficiency, one of the most important things you can do is to quit smoking, and protect yourself from secondhand smoke exposure. Tobacco smoke irritates and damages the lungs. For help with quitting smoking, see ATS Fact Sheets listed in Resources. In people with alpha-one antitrypsin deficiency, inhaling tobacco smoke speeds up the damage to the lungs.

There are a number of other steps you can take to reduce the risk of breathing problems if you have alpha-one antitrypsin deficiency. These include:

- Stay indoors when air quality is poor such as with an ozone pollution alert. Keep windows closed.
- Avoid dust whenever you can. If you have to clean, wear a mask, particularly when shaking rugs, vacuuming, sweeping, and dusting.
- Avoid any occupation where there is dust exposure (See ATS Patient information Series piece on Occupational Lung Disease at patients.thoracic.org).
- People with alpha-one antitrypsin deficiency may be more severely affected by respiratory infections. To reduce the risk of developing serious complications from these infections, get pneumococcal pneumonia vaccines and avoid people who are ill.
- Wash your hands frequently—it’s the best way to avoid catching a cold or the flu.
- Contact your health care provider at the first sign of a cold or other lung problem so you can try to keep it from getting worse.
- Exercise regularly to stay in shape.
- Avoid excessive alcohol—drinking may increase the risk of developing liver problems in people with alpha-one antitrypsin deficiency.
- Ask your doctor whether you would benefit from AAT augmentation therapy.


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Resources
American Thoracic Society
http://patients.thoracic.org/?page_id=283
(Sections on Pulmonary Rehabilitation, Tobacco, and COPD)
Alpha-1 Foundation
http://www.alphaone.org

Action Steps

- Quit smoking and stay away from second-hand smoke
- Exercise regularly to stay in shape
- Avoid exposure to outdoor and indoor pollution like ozone, dust and fumes
- Get your flu and pneumonia vaccines as recommended by your health care provider
- Contact your health care provider early if you have a cold or other respiratory illness
- Avoid excessive alcohol consumption
- Get regular health check-ups and lung function testing

Health Care Provider’s Contact Numbers/E-mail Address:

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