Partial Alpha 1 Antitrypsin Deficiency (Pi MZ)

An information leaflet for patients and families

Building healthier lives

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Introduction

This leaflet is written for people who are found to be carriers of alpha 1 antitrypsin deficiency (α1AD).

What is α1 antitrypsin (α1AT)?

α1AT is a chemical made in the liver that circulates in your bloodstream.

Why is α1AT important?

α1AT protects lung tissue from an enzyme (elastase) released by white blood cells. Elastase fights infection in the lungs. However if not tightly controlled by α1AT, elastase can attack healthy lung tissue.

What is α1AD?

Individuals with α1AD have very low levels of α1AT in their bloodstream. Carriers have levels which are low but much closer to normal.

How common is being a carrier of α1AD?

Around 1 in 10 people are carriers for the S or Z variant. 4% (1 in 25) of the Northern European population carry Z and 6% (1 in 17) carry S.

What are the effects of being a carrier of α1AD?

All the information in studies carried out throughout the world indicates that having partial deficiency (Pi MZ) is not associated with any specific tendency to develop severe health problems. There has been some information to suggest there may be a very slight increased tendency to developing liver cirrhosis. There may be a possible slight relationship to the development of asthma, but these people are certainly no more likely to develop emphysema than any other healthy person who smokes.
What causes α1AD?
Our ability to make α1AT is inherited through genes passed on by both parents. One of these genes is known as Protease Inhibitor (Pi). It is this gene that makes the α1AT.

How is it inherited?
We all have two Pi genes. We inherited one gene from our mother and the other gene from our father. When a couple has a child they each pass on one of their Pi genes.

What are the different genetic types?
There are more than 100 different variants of the Pi gene. Most variants result in normal levels of α1AT in the blood, but some result in reduced levels or no α1AT. The most common variants are called M, S and Z.

Most people have two copies of type M variant (written as PiMM) and have normal levels of α1AT in the bloodstream.

PiZZ
Type Z results in low levels of α1AT in the bloodstream. Someone with two Z variants (PiZZ) has α1AD

What do carriers of α1AT need to do to look after their health?
1. Avoid smoking and passive smoking. Smoking attracts white blood cells to the lungs and speeds up the development of lung disease. Other lung irritants e.g. dust particles and certain chemicals, should be avoided where possible.
2. Ask family members or partners to get tested for α1AD.
The reason for doing this is chance. Everyone has a 3 to 6 chance in
100 of marrying or partnering another person with a similar partial
deficiency, as it is quite common in the general population.

If we identify such a partner this has very important implications for
their present or future children.

For instance, if a Pi Z deficient patient marries a partially deficient (Pi
MZ) person, 1 in 2 of their children is likely to have severe deficiency
(Pi Z) as well as their parent (the patient we originally found).

If on the other hand a person with partial alpha 1 antitrypsin
deficiency (Pi MZ) marries another patient with partial deficiency, 1 in
4 of their children, again by chance, should develop severe deficiency
of the Pi Z type.

It is the detection of these children that is the most important factor.
By understanding that they have the deficiency, they can be advised
with the help of their parents never to take up smoking and, if so, it
is highly unlikely that they will develop severe emphysema. We know
that prompt treatment of all chest problems is important in retaining
the health of these children and also they require advice on which jobs
would or would not be suitable for someone with their tendency to
develop lung disease.

In other words, it is the aim of doing these tests to prevent ill health in
children and grandchildren, nephews and nieces before it really starts.

Will my children have α1AD if I am a carrier?

If your partner is not a carrier you will not have a child with α1AD. Your children will each have a 50% chance of being a carrier.

If your partner is also a carrier (Pi MZ) there will be three possible outcomes in each pregnancy.

1. 1 in 4 (25%) chance you will both pass on the Z variant so your child will have α1AD.

2. 1 in 2 (50%) chance one of you will pass the Z type and the other will pass the normal variant so the child will be a carrier of α1AD.

3. 1 in 4 (25%) chance neither of you will pass on the Z variant so your child will not be a carrier of α1AD.
Further information

**Alpha-1 UK Support Group**
36 Cecil Avenue, Lipson, Plymouth. PL4 8SG
Tel: 01752 225 573
Email: info@alpha1.org.uk
Web: www.alpha1.org.uk

**Alpha-1 Awareness UK**
Alpha-1 Awareness UK, PO Box 8294, Bakewell, DE45 9BF
Email: info@alpha1.uk
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Please use the space below to write down any questions you may have and bring this with you to your next appointment.
The Trust provides free monthly health talks on a variety of medical conditions and treatments. For more information visit www.uhb.nhs.uk/health-talks.htm or call 0121 371 4323.

*With thanks to the Clinical Genetics Unit, Birmingham Women’s NHS Foundation Trust

If you need more advice about alpha 1 antitrypsin deficiency please contact:

**Alpha-1 Specialist NHS Service/ADAPT Project**

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Website: www.uhb.nhs.uk/alpha-1-antitrypsin-deficiency.htm