The term Thalassemia is derived from the Greek word "Thalassa" meaning "the sea" as the patients were initially identified in the Mediterranean area. It is also termed as "Mediterranean Anaemia". It was first identified by Cooley & Lee in 1925, hence also named as Cooley’s Anaemia.

Thalassemia is an inherited blood disorder in which the body is unable to make adequate haemoglobin. Haemoglobin is present in the red cells. Normally red cells survive for 120 days but in Thalassemia red cell survival is reduced.

According to joint WHO-March of Dimes report 2006, approximately 7% of the global population is a carrier for Haemoglobin disorders with no health problems. Around 300,000 – 500,000 children are born annually with a severe haemoglobin disorder. Over 1,00,000 Thalassemia Major are born annually world over. Throughout the world there is a "Thalassemia Belt" that includes from countries around Mediterranean Sea like Italy, Greece, Cyprus, Sardinia and passes through West and Central Asian countries like Turkey, Saudi Arabia, Iran, Afghanistan onto Pakistan, India and South East Asian countries like Burma, Thailand & Indonesia,. Migrants/Descendants from these areas to other parts of world are also at high risk of carrying BTT.

**What is Thalassaemia?**

**Introduction**

Thalassaemia may have originated over 50,000 years ago. In a valley south of Italy and Greece now covered by the Mediterranean Sea. The name Thalassaemia is derived from a Greek word meaning sea. But Thalassemia was recognized as a clinical entity by Dr Thomas Cooley and Dr Pearl Lee who described five cases of Thalassaemia in 1925.

The thalassemias are a diverse group of genetic blood diseases characterized by absent or decreased production of normal hemoglobin, resulting in a microcytic anemia of varying degree. Thalassemia includes a number of different forms of anemia (red blood cell deficiency). The two main types are called alpha and beta thalassemias, depending on which part of an oxygen-carrying protein (called hemoglobin) is lacking in the red blood cells.

The alpha thalassemias are concentrated in Southeast Asia, Malaysia, and southern China. The beta thalassemias are seen primarily in the areas surrounding Mediterranean Sea, Africa and Southeast Asia. Due to global migration patterns, there has been an increase in the incidence of thalassemia in North America, primarily due to immigration from Southeast Asia.

**Alpha Thalassaemia**
The alpha thalassemias are caused by a decrease in production of alpha globin chains due to a deletion or mutation of one or more of the four alpha globin genes located on chromosome 16. The alpha thalassemias can be generally categorized as: Silent Carrier, Alpha Thalassemia Trait, Hemoglobin H disease, Hemoglobin H-Constant Spring, and Alpha Thalassemia major. Frequently, the diagnosis of alpha thalassemia trait in a parent is discovered after the birth of an affected child.

The most severe form of alpha thalassemia, results in fetal or newborn death. Most individuals with alpha thalassemia have milder forms of the disease, with varying degrees of anemia.

**Beta thalassemia**

Beta thalassemias, ranges from very severe to having no effect on health. There are three kinds of beta thalassaemia.

**Thalassemia major,** the most severe form, is also called Cooley's anemia, named after the doctor who first described it in 1925.

**Thalassemia intermedia** is a mild Cooley's anemia.

**Thalassemia minor** (also called thalassemia trait) may cause no symptoms, but changes in the blood do occur.

**Causes**

**General**

Thalassemia is an inherited condition. The disease is passed on through parents who carry the thalassemia gene in their cells. A "carrier" has one normal gene and one thalassemia gene also called as "thalassemia trait." Most carriers lead completely normal, healthy lives.

When two carriers become parents, there is a one-in-four chance that any child they have will inherit a thalassemia gene from each parent and have a severe form of the disease. There is a two-in-four chance that the child will inherit one of each kind of gene and become a carrier like its parents; and a one-in-four chance that the child will inherit two normal genes from its parents and be completely free of the disease or carrier state. These odds are the same for each pregnancy when both parents are carriers. The clinical severity of thalassemia varies tremendously depending on the exact nature of the genes that a person inherits.

Blood is made up of plasma (fluid), red blood cells, white blood cells and platelets. The white cells protect your body against and fight infection and the platelets are responsible for normal blood clotting. The red blood cells carry red blood protein called haemoglobin. Haemoglobin contains iron and transports oxygen from your lungs around
the body. Anaemia is caused by reduced haemoglobin. If the anaemia is mild it does no harm and may not be noticeable.

A normal haemoglobin molecule contains four protein (globin) chains (two alpha globin chains and two beta globin chains) Different genes are responsible for producing each chain. In thalassaemia there is an inherited defect in one of these genes. If the alpha chain is affected this causes alpha thalassaemia. If the beta chain is affected this causes beta thalassaemia.

**Causes of alpha thalassaemia**

In alpha thalassaemia there is a decrease in the production of alpha globin chains due to a deletion (missing) or mutation (abnormal change) of one or more of the four alpha globin genes located on chromosome 16.

**Causes of beta thalassaemia**

Beta thalassemias are caused by mutations on chromosome 11.

When two carriers become parents, there is a one-in-four chance that any child they have will inherit a thalassemia gene from each parent and have a severe form of the disease. There is a two-in-four chance that the child will inherit one of each kind of gene and become a carrier like its parents; and a one-in-four chance that the child will inherit two normal genes from its parents and be completely free of the disease or carrier state. These odds are the same for each pregnancy when both parents are carriers.
Symptoms

Alpha Thalassaemia

Alpha plus Thalassaemia

This is very common in some ethnic groups. There are no symptoms and it usually goes unnoticed. There may be slight iron deficiency (anaemia) if the blood is tested for some reason. Sometimes people may be mistakenly diagnosed as having iron deficiency anaemia and be treated with iron medications unnecessarily.

Alpha Zero Thalassaemia

There are no symptoms and you are perfectly healthy. However if both parents have alpha zero thalassaemia they have a 1 in 4 chance of having a baby who has alpha zero thalassaemia major which is incompatible with life; the baby is often born prematurely and is dead or dies shortly after birth.

Beta Thalassaemia

Beta Thalassaemia Trait (Carrier)

There are usually no symptoms and you are perfectly healthy, however there are an increased number of cells and they are smaller than those without the condition. It can cause mild anaemia because slightly less haemoglobin is produced than normal. This usually does not cause you any symptoms and cannot be treated by increased iron intake.

Beta Thalassaemia major

Between births and three to six months, the baby with Beta thalassaemia major will seem normal and quite healthy. The baby will then begin to show symptoms of anaemia (they become pale) there may be shortness of breath, jaundice and an enlarged spleen.

Without treatment your child’s bones will grow abnormally and death will occur early in childhood. Also if the condition is poorly treated or between transfusions your child will be pale, lethargic and breathless. There may be yellowing (jaundice) of the eyes and skin due to excessive breakdown of red blood cells. Also with poor treatment growth may be delayed, there may be osteoporosis of the bones and the spleen may be damaged.

Beta Thalassemia intermedia
Children with thalassemia intermedia may develop some of the same complications, although in most cases, the course of the disease is mild for the first two decades of life.

**Treatment**

Treatment for Beta thalassaemia major involves having regular blood transfusions, which take place every two to four weeks, depending on the severity of the anaemia. Excess iron builds up in the body from these regular transfusions. This is called iron overload. If this is left untreated, iron will build up in the body, leading to a condition called haemosiderosis. This can cause serious long-term damage such as heart failure and liver failure.

Iron overload is kept under control by treatment with medicines Desferrioxamine, Deferiprone and/or Defrasirox to achieve a more normal level of iron in the body. These medicines work by binding (the chemical term is “chelating”) with the iron in the blood and the chelated iron is then removed by the kidneys (excretion) from the body. Desferrioxamine has to be given by injection, usually by a slow injection under the skin via a small device or pump over eight to twelve hours. The Patient is taught how to do this yourself at home. The amount of desferrioxamine and how often it is given depends on the how much iron you have in your body i.e. the amount of iron overload.

As you will be receiving this medicine regularly (often daily), it can become tiresome for you or your career. So, it is important to understand why desferrioxamine doses should not be missed. Sticking to desferrioxamine treatment routines helps protect against serious complications in later life, such as diabetes and heart disease. There are also more immediate benefits such as prevention of nausea and sickness caused by iron overload.

Deferiprone is another medicine used to treat iron overload. It works in a similar way to desferrioxamine but is given by mouth. This medicine is only given to patients in whom desferrioxamine is not suitable or is not tolerated.

Like all medicines, both desferrioxamine and deferiprone may have side effects. You should refer to the manufacturer’s patient information leaflet for further information as well as talk to your doctor or pharmacist if you have any particular concerns. If you are receiving either of these medicines you need to be closely monitored by having regular blood tests and eye and ear examinations. Checks on body weight and height every three months may be carried out in children because their growth can be affected by the condition and, in some cases, by the medicines.

Beta thalassaemia intermedia treatment may be the same as for Beta major but as the anaemia is less severe, the need for transfusions or the frequency of transfusions and need for iron overload treatment will be different according to the severity of the anaemia.

**Supplementary treatment**
The time taken for iron to be removed from your body by desferrioxamine is improved by taking a daily dose of ascorbic acid (vitamin C). This is prescribed by your doctor according to your age and is usually started one month after starting desferrioxamine treatment. Ascorbic acid prescribed for this purpose should be taken separately from food because it also increases the absorption of iron in food, which you must avoid.

If you have heart problems, you will not be given ascorbic acid because its combined effect with the iron in the blood may make the heart problem worse. Any other multivitamin and mineral supplements should not be taken unless prescribed.

Other Treatments

Vene-section - therapeutic removal of blood from the body - to help deal with excess iron build-up

Surgery – removal of the spleen (splenectomy) is sometimes required if the spleen is damaged because it has to remove a larger number of abnormal red cells from the circulation.

Management and reduction of iron in the diet - usually only a small amount of iron is absorbed from the diet, however the absorption is increased when haemoglobin in the blood is low, as in beta Thalassaemia major. This is especially true between transfusions. Therefore iron in the diet should be low (red meat is especially high in iron). You should be referred to a dietician for advice on diet.

Bone marrow transplantation – this can provide a cure for Beta thalassaemia major. Bone marrow is transplanted from a matched unaffected sibling or unrelated donor. It is best done when the child is very young. However, the procedure is painful and, although success rates are improving, they are unpredictable.

There are many emotional, psychological and social effects for the person with thalassaemia and their family, particularly as self-management is so important. Psychological support is important in managing chelation therapy and other aspects of the condition. You may find getting involved with local support groups and voluntary organizations helpful.

FAQ’s

1) What are the forms of Thalassaemia?
There are two forms of Thalassaemia.
Thalassaemia Major
Thalassaemia Minor (Trait)
2) What is Thalassaemia Major?
Thalassaemia major is sometimes known as Cooley’s Anaemia, Homozygous, Bête Thalassaemia or Mediterranean Anaemia. Is serious inherited childhood anaemia. Children with Thalassaemia major cannot make enough haemoglobin. Because of this, their bone marrow cannot produce enough red blood cells. The red blood cells that are produced are nearly empty. (Thalassaemia newsletter Sep 1988 page 1)

3) What is Thalassaemia Minor?
People with Thalassaemia Minor, sometimes known as Trait, carry Thalassaemia but they are not ill. They are completely healthy and normal but some of them have slight anaemia. Most people with Thalassaemia Minor do not even know that they have it. It is only discovered if the person has a special blood test or if they have a child with Thalassaemia Major. It is important to know if you have Thalassaemia Minor later in life. The reason for this is that it may cause some problems if the person and their partner want to start a family. Thalassaemia minor’s red blood cell are also different from normal blood cells. (Thalassaemia Newsletter July 1988 page 1)

4) Who are Carriers of Thalassaemia and what is the number of people who are carriers of Thalassaemia around the World?
There are 100,000 children born in the world with Thalassaemia major. In Australia there are about 300 people with Thalassaemia. In Victoria there are about 180 people. There may also be 60,000 people to 100,000 people in Victoria who carry Thalassaemia Minor but don't even know it. (Hall 1994)

5) Who is likely to carry Thalassaemia?
People who are likely to carry the gene of Thalassaemia are people with Mediterranean descent, for example Cyprus, Egypt, Greece, India, Italy, Lebanon, Malta, Middle East, Turkey and some parts of South East Asia.

6) Can Thalassaemia major patients also carry other illnesses?
Thalassaemia major patients can also carry other illnesses such as Sickle Cells, Diabetes, liver dis.-function, and other illness that non Thalassaemia people can get for example Cancer.

7) What are known cause for Thalassemia?
There is not a known cause for Thalassaemia except that is inherited through the genes.

8) What is the "quality of life" for a Thalassemic?
A chronic illness always causes some limitation of quality of life, especially when it requires frequent and complex treatment, as Thalassaemia does. The treatment should not interfere with a Thalassemics life. In particular doctors and hospitals should make the effort to arrange out-patient visits and visits for transfusions so they interfere as little as possible with normal life. Treatment should not interrupt schooling or work.
9) How long can a person with Thalassemia major live?
These days most Thalassemics grow up to become adults, and earn their own living. Most also find a partner and get married. Now a number of Thalassaemia major patients have their own children.

It is very hard to know the answer for Thalasseemics who are well at present. The disorder and its influence are changing almost from day to day, because of advances in treatment. Thalassemic patients are now living longer. Today it is reasonable to think that people with Thalassaemia major, who have been well treated from the beginning, may well live as long as people without Thalassaemia. Only time will tell. Even so Thalassemics live with more risks than non Thalassemic, because of the amount of medication and treatment they receive. But all medical treatments include some risk.

10) Can people with Thalassaemia major and minor have healthy children?
People with Thalassaemia major can have babies only if their partner does not carry any sort of Thalassemia. But all Thalassaemia major's patients children will carry Thalassaemia minor.

If a Thalassaemia major partner does not carry any Thalassaemia gene none of the children would have Thalassaemia major.

The chance of having a baby with Thalassaemia minor decrease if their partners have Thalassaemia minor.

11) History of treatment and Victoria treatment
In the 1940's, blood transfusions were introduced to treat Thalassaemia major patients on a rare bases due to a concern about transfusion iron overload causing organ damage or dysfunction and ultimate fatality. These transfusions had the effect of raising the haemoglobin levels from three to eight grams and therefore increasing the life expectancy between sixteen to eighteen years.
In the late 1960s, high transfusion on average high transfusion regimens were introduced involving regular blood transfusions on average every 4-8 weeks, depending upon individual requirements. The aim of this regimen is to maintain haemoglobin levels as close as possible to the normal range. Desferrioxamine was given by a daily intramuscular injection and was effective for up to 6 hours following each injection. Occasionally there were side-effects associated with this method, example hypotension, nausea and vomiting.

12) Do Thalassaemia person need to be on a special diet?
Thalassaemia major patents should try to keep away from high in iron foods such as red meat, liver, kidney, green leafy vegetables such as spinach, some breakfast cereals, whole meal breads and alcohol. Although this is recommend, patients do not have to stick with this diet.
12) Is the treatment effective?  
This treatment is very successful and most children treated with blood transfusions and Desferal can now lead fairly normal healthy lives.

13) How is the treatment improving?  
Treatment today is more advanced than what it was. A Thalassaemia person can enjoy a good quality life, lively in normal activities such as sport, study, work, marriage and family.

An oral version in pill or liquid form, would greatly improve the quality and productivity of the lives of Thalassaemia major patients would be better. There have been scientists working on this pill it is known as L1. The results from the L1 have been good except of one specific side affect called "Neutropenia" which reduced the ability of the body to cope with infections so the Thalassemic person has a risk to infections and can die due to this. But Ciba Geigy (company which researches, and will make the drug when it is discovered) promised to be more involve on the oral chelators. Ciba Geigy believes the oral chelator will be available before 1996 to 1997. Also scientists believe that a cure for Thalassaemia is possible through gene therapy and bone marrow transplantation. Gene therapy would correct the defected gene in Thalassaemia patients. This is done by transferring normal gene into the patients own bone marrow cells. But A bone marrow transplant proven to be a success however the odds against success with possible death over failure.