This fact sheet describes the chromosome condition trisomy 13 and includes the symptoms, cause and any treatment or testing which is available.

**In summary**
- Trisomy 13 is a chromosome condition also known as Patau syndrome
- Babies with trisomy 13 usually have distinctive features, severe intellectual disability and other physical developmental problems
- Trisomy 13 is caused by having an extra copy of chromosome number 13.

**WHAT IS TRISOMY 13?**
Trisomy 13 is also known as Patau syndrome. It is a condition which is considered very serious and most babies with trisomy 13 do not survive to birth.

The combination of features present in babies affected with trisomy 13 can lead to many different developmental problems.

Some general symptoms include:
- Developmental delays, profound intellectual disability, unusually small eyes, an abnormal groove or split in the upper lip (cleft lip), an incomplete closure of the roof of the mouth (cleft palate), undescended testes in boys, and the presence of extra fingers and toes
- Incomplete development of certain parts of the brain (e.g. the forebrain), kidney abnormalities, structural heart defects at birth such as an abnormal opening in the partition dividing the upper or lower chambers of the heart.
- A relatively small head with a sloping forehead, a broad, flat nose, widely set eyes, vertical skin folds covering the eyes inner corners, scalp defects and low-set ears
- Failure to grow and gain weight at the expected rate and severe feeding difficulties, diminished muscle tone and episodes in which there is temporary cessation of spontaneous breathing
- Life-threatening complications may develop before birth, during infancy or early childhood.

**WHAT CAUSES TRISOMY 13?**
As mentioned above, trisomy 13 is caused by the presence of an extra full (or partial) copy of chromosome number 13.

In each human cell, except the egg and sperm cells, there are 46 chromosomes. Chromosomes are found in pairs and each pair varies in size.

Thus there are 23 pairs of chromosomes, one of each pair being inherited from each parent.
- There are 22 numbered chromosomes from the largest to the smallest: i.e. 1-22. These are called autosomes
- There are also two sex chromosomes, called X and Y.

In females, cells in the body have 46 chromosomes (44 autosomes plus two copies of the X chromosome). They are said to have a 46,XX karyotype. Eggs (female reproductive cells) are different as they only contain half of the chromosomes (23 made up of 22 numbered chromosomes and an X chromosome).

In males, cells in the body have 46 chromosomes (44 autosomes plus an X and a Y chromosome). They are said to have a 46,XY karyotype. Sperm (male reproductive cells) are different as they only contain half of the chromosomes (23 made up of 22 numbered chromosomes and an X chromosome or a Y chromosome).
When egg and sperm cells are formed, the chromosome pairs separate so that there is only one of each pair in these cells i.e. 23 chromosomes instead of 46. A baby is conceived when the egg from the mother and the sperm from the father come together.

The baby would then have two copies of each chromosome (46 chromosomes in total) just like the parents. One copy of each chromosome would have come from the mother and one copy from the father (see Figure 37.1).

Figure 37.2 shows a chromosome picture (karyotype) from a male where each chromosome has been numbered from the largest (chromosome number 1) to the smallest (chromosome number 22) and arranged in pairs in order of size.

We know these chromosomes are from a male because of the X and a Y.

Sometimes, when the egg and sperm are forming, a mistake can occur so that the chromosome pairs do not separate in the usual way. The result is an egg or sperm cell that has only 22 chromosomes while others have 24 chromosomes.

If an egg or sperm carrying 24 chromosomes combines with an egg or sperm carrying the usual 23 chromosomes, the result would be an individual with cells in which there are 47 chromosomes instead of the usual 46 (See Figure 37.3).

- There would be three copies of a particular chromosome in the cells rather than two. This is called trisomy.

The chromosome pattern in people with trisomy 13 usually includes an extra copy of chromosome number 13. This extra chromosome causes an imbalance and is the reason why there are differences in people with 3 copies of chromosome 13 material and those with the usual 2 copies of chromosome 13.

The presence of the extra copy of chromosome 13 causes the intellectual and physical characteristics of trisomy 13. Individuals with this condition usually have three copies of chromosome number 13, i.e. 47 chromosomes in their cells instead of 46. Trisomy means three bodies.

Figure 37.3 When the egg has 24 chromosomes, and the sperm has the usual 23, the baby’s cells will contain 47 chromosomes instead of 46.
**Figure 37.4**: Chromosome picture (karyotype) from a female with trisomy 13. In this cell, there are 47 chromosomes including three copies of chromosome 13 instead of the usual two.

Figure 37.4 is a picture (karyotype) of the chromosomes from a female with trisomy 13.

**HOW IS TRISOMY 13 INHERITED?**
In most cases where trisomy 13 is caused by a complete extra copy of chromosome 13, that person will be the first and only person affected by the condition in that family. It is usually assumed that if the parents of a person with trisomy 13 have the usual two copies of chromosome 13, then the extra 13 in their child was a result of an egg or sperm with 24 instead of 23 chromosomes being fertilised.

Older mothers have a higher chance of having these errors.

**IS THERE ANY TESTING AVAILABLE FOR TRISOMY 13?**
Chromosome testing of a baby who is suspected of having trisomy 13 can be done using a blood sample. A doctor may suspect a baby has the condition based on observing the characteristic features or symptoms associated with the syndrome.

**Prenatal testing and PGD**
Testing for trisomy 13 may be offered during pregnancy to some couples, especially for those women who are 35 years or older.

Prenatal tests can occur in a number of different ways and at certain stages of the pregnancy. In general, there are two main types of prenatal test – a screening test and a diagnostic test.

*Screening tests* give a risk or estimate of the probability that a baby has a health problem. These tests do not generally look directly at a sample from the developing baby and are therefore very safe. Included in the group of screening tests are ultrasounds, non-invasive prenatal testing, first trimester screening and second trimester screening.

*Diagnostic tests* provide a more accurate result since they are directly testing the baby. Because of this, in a very small number of cases, the test may also cause a miscarriage. Included in the group of diagnostic tests are ultrasounds, chorionic villus sampling (CVS) and amniocentesis.

Testing during pregnancy is optional and should be discussed in full with your doctor, midwife or genetic counsellor. Making a decision to have a test or not is always up to you.

It may also be possible to undergo pre-implantation genetic diagnosis (PGD) screening for trisomy 13 on an embryo created using in vitro fertilisation (IVF). These options are best discussed and considered before pregnancy, when possible, in order to ensure all possible risks, benefits and outcomes can be explored.