Introduction

The purpose of this booklet is to provide families with information about prenatal testing results that indicate a change in the sex chromosomes. This can be a very anxious time for families, and often little information is available to people who are not in the field of genetics. This booklet will provide you with some information, but it probably will not answer all of your questions. Please talk to a genetic counselor or a medical geneticist for more specific information about your situation. They are there to help. This booklet may also help you explain the test results to your family and health care providers.
I. Cells

The testing that was performed on your fetus was done by examining cells from either the amniotic fluid (if you had amniocentesis) or the placenta (if you had chorionic villus sampling). When the cells were studied, the packages of genetic information, called chromosomes, were examined. Your body, and your baby’s body are made up of many billions of cells. Each cell contains a complete set of chromosomes. The cells of the fetus, placenta, and amniotic fluid, all came from the fertilized egg. Therefore, the chromosomes studied from amniotic fluid or the placenta match the chromosomes of the fetus.

This is an illustration of a cell. It is so small that it can only be seen using a microscope:

II. Chromosomes

There are usually 46 chromosomes in each cell. They are like a set of cookbooks. Each chromosome or cookbook contains thousands of recipes that are pieces of information or instructions. These instructions or recipes are called “genes.” Therefore, the chromosomes are packages of genes, which direct the body’s development. For instance, there are genes that tell whether a person will have blue eyes or brown eyes. All of the information that the body needs to work is included in the chromosomes. The chromosomes contain the blueprint for growth and development. Scattered over the 23 pairs of chromosomes are about 30,000 genes. Even a very small piece of a chromosome contains many different genes. The precise location - or even the exact number - of all the genes is not known. Chromosome studies do not include a detailed examination of each gene.

Chromosomes come in pairs. One member of each pair comes from the father’s sperm cell and the other comes from the mother’s egg cell. In other words, the baby receives half its genetic material from the mother and half from the father. When the chromosomes are examined in the laboratory under the microscope, they look like this:

III. The Karyotype

In order to study the chromosomes, the cells obtained from amniocentesis or chorionic villus sampling are prepared in the laboratory so they can be seen under the microscope. The cells are photographed, the chromosomes are cut out of the photograph, and lined up by their sizes and characteristic light and dark banding patterns. Usually, a boy has an X and a Y chromosome and a girl has two X chromosomes. This picture is called a “karyotype.”

This picture shows a normal male karyotype:

This picture shows a normal female karyotype
IV. Differences in Sex Chromosomes

Although most people have chromosomes that look like these photographs, some people have a different number of sex chromosomes. Others have a sex chromosome with an unusual structure. Most people with changes in their sex chromosomes are healthy, and look like other members of their families. Sex chromosome abnormalities are rarely diagnosed at birth. Unless the mother has an amniocentesis or chorionic villus sampling, it may be many years before the chromosome change is diagnosed. Sometimes it is never diagnosed. A sex chromosome change may be discovered when a child does not go through puberty normally, or much later, has trouble having children.

The most common reasons for studying a fetus’s chromosomes are: the mother’s age, a positive maternal serum screening test result, or an abnormal ultrasound evaluation. In these situations, the laboratory is mainly looking to make sure the baby does not have an extra “autosome.” “Autosomes” are the chromosomes other than the sex chromosomes. As an example, babies with Down syndrome (trisomy 21) have an extra chromosome 21 in their cells. Since all of the fetus’s chromosomes are examined during a prenatal chromosome study, changes in the sex chromosomes are also detected.

There are several different types of sex chromosome changes: an extra or missing copy of the entire X or Y chromosome, an extra or missing part of the X or Y chromosome, or a rearranged X or Y chromosome. Changes in the number of chromosomes result from an error in the formation of the egg or sperm cell. When this happens, the other cells in the parents’ bodies are normal. Neither parent needs to have a chromosome study of their own cells. An extra, missing, or rearranged sex chromosome may be inherited from a parent, or have occurred for the first time during the formation of the egg or sperm. If a rearranged chromosome is discovered, your geneticist may recommend testing the parents’ chromosomes.

Parents often feel guilty when their fetus has a chromosome change. However, nothing either parent did “wrong” caused the sex chromosome change.

V. What can these results mean for your fetus?

Most babies with changes in their sex chromosomes are healthy at birth, without serious birth defects. The sex chromosome change may not cause any obvious problems in your baby’s early growth and development. Some people with sex chromosome changes never even learn they have a sex chromosome abnormality. But, there is a chance that there will be problems. Learning difficulties and emotional problems are more common in people with a change in their sex chromosomes. These problems cannot be identified prenatally or early in life.

Other problems can occur, and depend on which chromosome change is present. Your physician or genetic counselor may recommend additional studies of your fetus. Since it is not possible to identify all birth defects before any baby is born, there will still be a risk for physical problems even if all the follow up studies are normal.

Parents often wonder if a change in the sex chromosomes means that the baby will be born with both male and female sex organs (ambiguous genitalia), or that their child will be homosexual. Having both male and female sex organs or being homosexual is no more likely in a child with a change in the sex chromosomes than in someone with the typical sex chromosomes.

For some people, the additional risk for problems in the fetus is too high, and they choose to end the pregnancy. Others choose to continue the pregnancy. The decision whether to stop a pregnancy is a very personal one. It is usually a very difficult decision, and may take time to make. Each family must consider their own feelings and the risks. Your genetic counselor or health care provider is available to help you make the best decision for your family. Let them know how they can help.

### Klinefelter Syndrome

You have recently learned that your fetus has Klinefelter syndrome. You had probably never heard of this condition before you had prenatal diagnosis. You may now be making important decisions about how to proceed. This booklet was created to answer some of the many questions you may have.

### What is Klinefelter syndrome?

Klinefelter syndrome is the name given to a set of physical features by a doctor, Harry Klinefelter, MD. He named the condition before the underlying cause was known. We now know that the features observed by Dr. Klinefelter are caused by a change in the sex chromosomes. Approximately 1 in 500 boys and men has Klinefelter syndrome.
What causes Klinefelter syndrome?

Boys and men who have Klinefelter syndrome have one Y chromosome and two X chromosomes instead of the usual one of each. This chromosome change is written as “47,XXY.” There are 47 chromosomes instead of the usual 46, and there are two X chromosomes instead of the usual one X and one Y (in boys) or two X’s (in girls). The extra X chromosome was gained either during the formation of the sperm or egg that later came together to form the fetus, or during early fetal development soon after conception. The extra chromosome can never be removed. Klinefelter syndrome happens by chance. Nothing the parents did caused it to happen, nor could they have done anything to prevent it from happening.

This is a karyotype from a man with Klinefelter syndrome:

![Karyotype of 47,XXY](image)

What physical features are to be expected with Klinefelter syndrome?

The features of Klinefelter syndrome are variable from one person to another. Some individuals with Klinefelter syndrome have such mild features that they are never diagnosed. Newborns with Klinefelter syndrome look like other newborn boys. In childhood, these boys are often taller than other boys, and this continues into adulthood. Boys with Klinefelter syndrome usually enter into puberty normally. However, because of the extra X chromosome, the testicles do not develop completely and therefore do not make as much testosterone as they should. Boys and men with Klinefelter syndrome may be less muscular than other males in their families and may not be as coordinated. Some breast development may occur and they may develop less facial and body hair. Men with Klinefelter syndrome have normal sexual function but are usually unable to produce sperm and are therefore unable to father children (infertile).

Each feature seen in Klinefelter syndrome can also be seen in boys and men with the usual number of chromosomes.

What mental or social features are to be expected with Klinefelter syndrome?

Mental retardation is not usually a feature of Klinefelter syndrome. Most men and boys with Klinefelter syndrome have normal intelligence, but they may have slightly lower IQs than other members of their families. An increased chance for speech and language problems can lead to delays in social skills and learning. Therefore they may need additional help to succeed in school. They are usually able to complete high school but may need to repeat a grade or receive special education. Some men with this condition have successfully gone on to college and graduate school.

Is there any cure for Klinefelter syndrome?

The sex chromosome change that causes Klinefelter syndrome can never be repaired. However, some therapies are available. Testosterone can be given at the time of puberty to help boys with Klinefelter syndrome develop a more masculine appearance. Any excess breast tissue that still develops can be surgically removed, if desired. The infertility cannot be cured but families can make a point of discussing alternatives to biological parenthood, such as adoption or the use of a sperm donor. A supportive home environment may reduce learning and behavioral problems.

Should I expect any complications during the rest of my pregnancy?

No. You have no higher chance of complications than the next person. The delivery and newborn period should be normal as well.
Will this happen again in future pregnancies?

Probably not. Your chance of having another baby with Klinefelter syndrome (or any other chromosome abnormality) is unlikely to be much greater than your age-related risk. Your other children, brothers and sisters, and other family members do not have an increased chance to have a child with Klinefelter syndrome or any other chromosome abnormality. However, once a woman has a baby with a chromosome abnormality, she is often worried in the next pregnancy. Prenatal diagnosis will be available in any of your future pregnancies.

Other questions you may have for your genetic counselor or health care provider:

- How do I know the prenatal diagnosis results are accurate?
- What are my options in this pregnancy?

Ask them about anything that concerns you.

Are there support groups available?

Yes, and talking with others who have “been there” can be very helpful. Try:

AXYS – Association for X and Y chromosome variations
Web: www.genetic.org
Email: info@genetic.org

Children with sex chromosome problems are frequently followed through a genetics clinic or by a pediatric endocrinologist. Call one of the phone numbers listed below to locate a clinic in your area.

In the Pacific Northwest

There are genetics clinics and prenatal diagnosis clinics in Alaska, Idaho, Oregon, and Washington.

Anyone can call for more information. To find the clinic nearest you, call:

### Alaska:
907-269-3430
http://dhss.alaska.gov/dph

### Idaho:
208-381-7339
http://www.healthandwelfare.idaho.gov

### Oregon:
971-673-0273
http://public.health.oregon.gov

### Washington:
253-395-6742
http://www.doh.wa.gov

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Copies of this booklet can be obtained from the Washington State Department of Health - Health Education Resource Exchange (H.E.R.E.) website:

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