STAGE 2 :
The non-invasive prenatal genomic test OR the diagnostic test
NON-INVASIVE PRENATAL GENOMIC TEST The genomic test is a screening test that will be proposed if you have a high probability of having
a baby with trisomy 21 (or trisomy 18) following the program's biochemical test. The genomic test is performed by taking a sample of the mother's blood and analyzing DNA fragments from the placenta. The genomic test screens for trisomy 21, trisomy 18 and trisomy 13
The genomic test is proposed because it is reliable and safe (there is no risk of miscarriage compared with the diagnostic test).

The genomic test might be proposed to you right away (instead of the biochemical test) if you are in one of the following situations:
you have had a previous pregnancy in which the baby had trisomy 21 , trisomy 18 or trisomy 13; you will be 40 years old or older at the time of delivery;
you are pregnant with twins (two babies); the test is prescribed following a consultation in genetic medicine.

RESULTS OF THE GENOMIC TEST
LOW PROBABILITY: this result indicates that the baby has a very low probability of having one of the test
trisomies, even if the result of the biochemical test showed a high probability initially. A "low probability" result of the genomic test is very reliable (over 99\%) and there is no need to conduct further investigations.

HIGH PROBABILITY: this result indicates that the baby is likely to have one of the three tested the baby is likely to have one of the three tested
trisomies. However, a "high probability" result is not $100 \%$ certain and only a diagnostic test can determine with great certainty whether the baby has one of the trisomies.

## DIAGNOSTIC TEST

Diagnostic testing, based on amniocentesis or biochemical test indicates a high probability.

The diagnostic test determines the number of each chromosome. It is reliable in telling whether or not techniques used to collect the sample for analysis present risks for the pregnancy, such as miscarriage.

RESULTS OF THE DIAGNOSTIC TEST
The baby does not have one of the tested trisomies (trisomy 21, trisomy 18 or trisomy 13.
The baby has one of the tested trisomies. This result is very reliable and allows you to make
a decision about your pregnancy

You can consult the table, which compares the genomic test and the diagnostic test, and discuss care professional.

PARTICIPATION
IN PRENATAL SCREENING
The decision to participate is entirely up to you. Some women want to have this information during the pregnancy and others dont.
Keep in mind that at any time and at any stage,
you can choose to end your participation
in the program
QUÉBEC PRENATAL SCREENING PROGRAM

For more information on prenatal screening and the limitations of biochemical and genomic tests, talk to the health care professional mo
or visit Québec.ca/depistage
Participating in the program might cause you stress Participating in the program might cause you stress
and anxiety before getting the tests, while awaiting and anxiety before getting the tests, while awaiting
your results or when receiving your results. You can discuss the decision on whether to participate in the Québec Prenatal Screening Program with your partne your loved ones, your doctor, your midwife, another health professional or other groups for parents who have a child with a trisomy.

Québec

## SUMMARY

## TRISOMY 21

Trisomy 21, also known as Down Syndrome, is one of the mos
1 in 800 babies).
A person with trisomy 21 has a $3^{\text {dr copy }}$
of chromosome 21. Therefore, the individual
has 47 chromosomes instead of 46 .
Trisomy 21 affects the child's intellectual
development, and that may vary from one child
to the next.
A child with trisomy 21 may suffer from other health
problems (e.g., heart defects).

## TRISOMY 18

Trisomy 18 is a rare trisomy (affecting approximately 1 in 4,500 babies at birth).
Trisomy 18 is caused by the presence of a $3^{\text {rd }}$ copy
of chromosome 18.
Trisomy 18 is often associated with severe birth defects.

Most pregnancies with a baby having trisomy 18 nd in miscarriage
Most babies born with trisomy 18 die soon after arth as a sult ofere heart and brain defect or respiratory problems.

## TRISOMY 13

Trisomy 13 is a rare chromosomal abnormality (affecting approximately 1 in 7,000 babies at birth). Trisomy 13 is caused by the presence of a $3^{\text {rd }}$ copy of chromosome 13.
isomy 13 is very severe and often associated with miscarriage or multiple defects.
born with trisomy 13 die soa after birth.

Trisomy 21, like trisomy 18 and trisomy 13 is not hereditary in more than $95 \%$ of cases and therefore occurs randomly. Less than 5\% of cases are inherited

The level of stimulation and support offered to these children impacts their development. In most cases, ndividuals with trisomy 21 will need varying levels of support throughout their lives. However, given a fair hance, individuals with trisomy 21 have the resources nd potential to develop deep emotional relationships oved ones.

## DESCRIPTION

OF THE QUÉBEC PRENATAL SCREENING PROGRAM

The program screens for trisomy 21, but it could also reveal trisomy 18 or trisomy 13 during the pregnancy. Conducting prenatal screening allows us to calculate the probability that your child has one of these trisomies. There are other chromosomal abnormalities but the public program, which is based on Canadian recommendations, does not screen for a decision on whether to continue the pregnancy tis up to mot It is up to you to decide whether to proceed with to make an informed decision. The choice is yours to make an informed decision. The choice is yours

The biochemical test
(with or without nuchal translucency)
The biochemical screening test reveals whether the probability of trisomy 21 is low or high. The test It could be combined or not with the nuchal translucency measurement

## BLOOD TEST

The biochemical test provides an analysis of the baby's eins or hormones present in the mother's blood.
 fist trimester of pregnancy, between the $10^{\text {th }}$ and $13^{\text {th }}$ weeks inclusively.
NOTE: It is important to take the blood test at the right time. A screening test taken in the second trimester provides results later in the pregnancy.

Any woman may carry a baby with trisomy 21 trisomy 18 or trisomy 13
The probability of having a child with trisomy 21, trisomy 18 or trisomy 13 increase with the pregnant woman's age. The younge

## IRST TRIMESTER ULTRASOUND

## first trimester ultrasound will be proposed betwee

 weeks 11 and 13 inclusively to see how your pregnan is progressing and identify possible abnormalities your baby. This ultrasound is used to precisely estimate how long you've been pregnant to allow a more reliable calculation of the probability of aving a baby with a trisomy. This ultrasound will also determine if there is a single baby or more than one The nuchal translucency measurement by ultrasound might also be proposed. It is performed between weeks 11 and 13 inclusively. It measures the thicknes of the fluid build-up at the back of the baby's neck. n babies with trisomy 21 , the fluid build-up is often thicker than normal. When available, the nuchal translucency measurement is combined with the trisomy Nuchal translucency measument could involve costs when performed in a private clinic
## ESULTS OF THE BIOCHEMICAL TEST

 WITH OR WITHOUT NUCHAL TRANSLUCENCY)LOW PROBABILTY: the probability that you re carrying a child with trisomy 21 is low and no additional tests are necessary.
A low probability does not guarantee that the baby does not have trisomy 21. Considering the limitations of the test, it does not allow all babies with trisomy 21 to be identified.
HIGH PROBABILITY: it is possible that you are tarrying a child with trisomy 21. The non-invasive prenatal genomic test or the diagnostic test will be proposed
A high probability does not necessarily mean that the baby has trisomy 21.
this stage, the test may reveal a high probability f trisomy 18.

