



Do you want to understand your genetic risk and want to learn about genetic testing?

Do you have family history or past medical history of a genetic disorder?

Do you find yourself lost in a Diagnostic Odyssey?

- Delayed diagnosis.
- Difficult access to technologies and expertise.
- Elevated cost.
- Lack of information and difficulty understanding.

Igenomix[®]
WITH SCIENCE ON YOUR SIDE

www.igenomix.com



GPDx

Genomic Precision
Diagnostic
by Igenomix[®]

The answers
you need in a
drop of blood

Your genetics matter,
learn about them



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V. 2021



Prevention and accurate diagnosis of genetic conditions must be a Global Health Priority

- Igenomix provides expertise, support and the highest technologies to give the best response and diagnosis to you patient.
- The highest diagnostic precision is obtained when appropriate technology is used.
- Igenomix's Genetic Specialists provide support to select the best technology according to clinical and family information.

What are the benefits of genetic testing?

- For a lot of disorders this is the **only way to make an accurate diagnosis** and help avoid additional unnecessary clinical investigations.
- Can **guide the clinician** in choosing the most suitable therapy and support for the patient.
- Reduces the uncertainty in a differential diagnosis and provides a directed focus for the physician **to provide appropriate treatment.**
- **Decreases referrals from specialist to specialist**, allowing a high quality patient care.

Our Services



Preconceptional

- Identify risk of carrying a genetic mutation before pregnancy.
- Decrease risk of genetic diseases.
- Genetic counselling.



Prenatal

- Identify high risk pregnancies.
- Use of technologies of genetic analysis of the fetal sample.



Neonatal

- Identify any variants that could cause disease in newborns.
- Prevent complications and increase life expectancy.

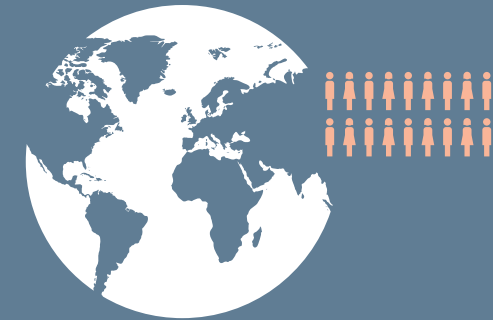


Childhood/Adulthood

- Identifies or rules out the mutation that causes the symptoms, confirming the diagnosis.
- Increases diagnostic yield for to ensure a high quality patient care.
- Identify risk of cancer and heart disease.

What are Rare Diseases?

Rare disease are a diverse group of conditions; very few people are affected by them in comparison to more common conditions like diabetes or heart disease.



Many people worldwide will be affected by a rare condition at some point in their lives.



~80%

Of rare diseases have a genetic origin.



~75%

Of rare diseases affect children.