

GeNSCREEN CARRIER SCREENING GENETIC TEST



**Pioneers in Reproductive Genomics
since 2011**

Screen Yourself and Safeguard Your Future Generation



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GeNSCREEN

Carrier Screening Genetic Test

Reproductive carrier screening either prior to conception or during pregnancy can help to identify your chance to have a child with serious genetic condition.

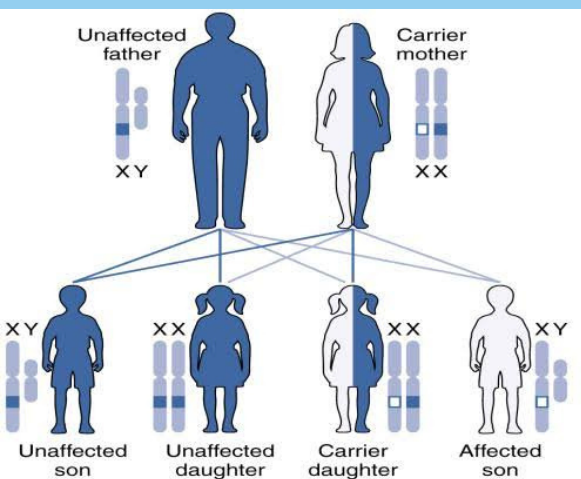
This test helps you to make decision about your reproductive options and prenatal care.



Your Path to a Healthier Future

Genetics is the 21st century 'Kundali', Traditional kundali in astrology is based on the positions of celestial bodies at the time of a person's birth, genetics delves into the code of life—DNA—offering a scientific view of inherited traits, predispositions, and even certain behavioral tendencies.

Genetics, much like astrology, holds a certain amount of predictive power, but it's grounded in biology and the environment rather than cosmic forces. It's like the blueprint of who we are, with the potential to reveal a lot about our future health, abilities, and even susceptibility to specific conditions.



Who are carrier

Carrier is a person who has a genetic change or a mutation or a variant in their DNA but in majority of the cases they are asymptomatic and are not having any associated health problems. Carriers can pass this mutation to their children and who may then develop a genetic condition with mild to severe symptoms

Best Test to select when

- Couple is having BOH
- Karyotype is normal.
- Karyotype/ Chromosomal Microarray is normal from POC.
- Family history of genetic disease.
- Consanguineous marriage
- Considering egg/sperm donor for IVF

Why GeNeCaRe

- Cutting edge technology
- Comprehensive analysis of all OMIM defined recessive disorders
- Screens over 4000+ genetic diseases
- Additional analysis for Spinal Muscular atrophy, Fragile X syndrome
- 3500+ Genes
- >95 coverage in coding regions of clinically relevant genes
- Enables the discovery of rare and novel mutations
- Pre and Post-test genetic counseling