

Genetic Service Solution for your Patients



Make
Advanced
Clinical
Decisions

Save &
Improve
More Lives

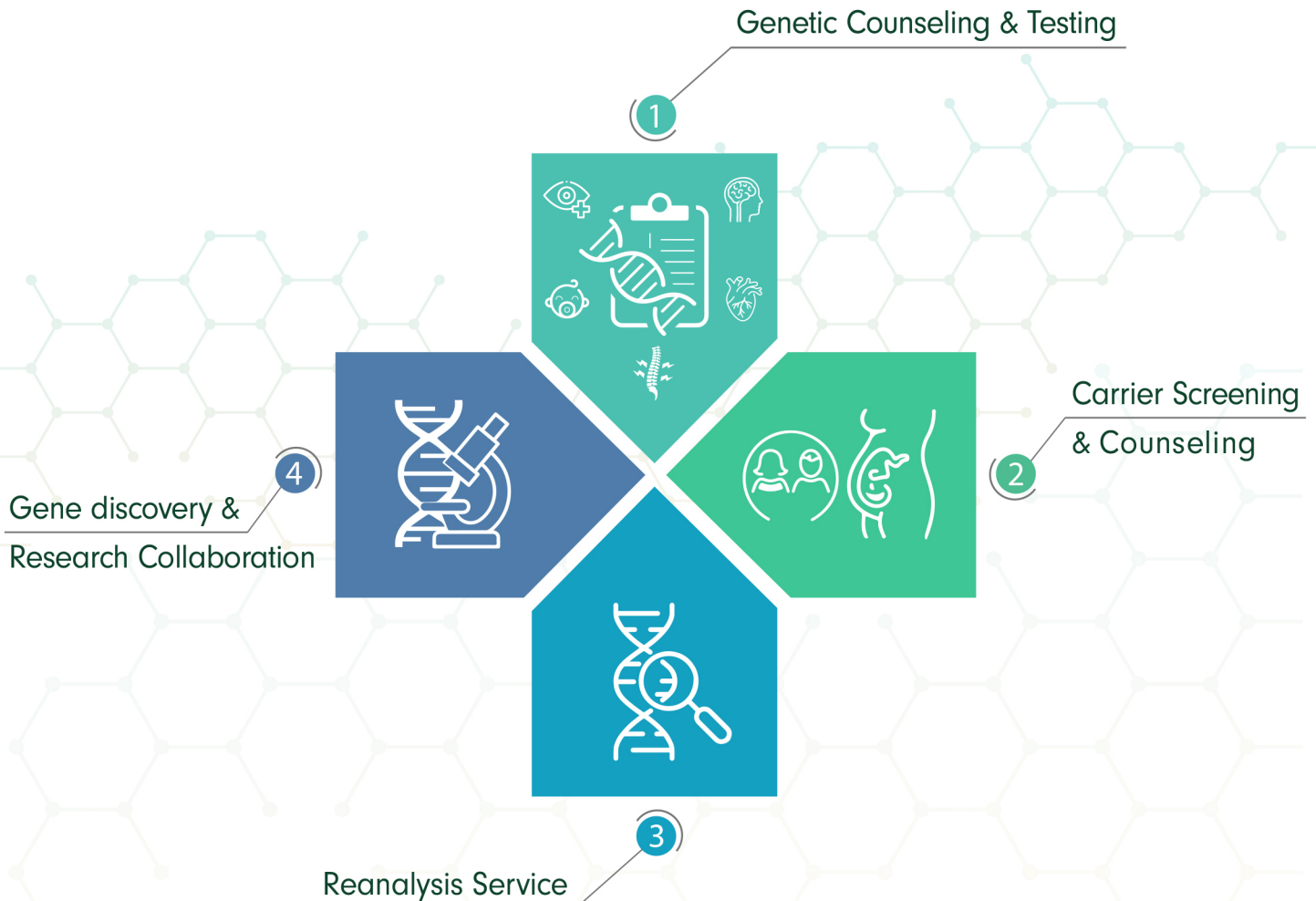
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Rare Disease Service Menu



1 In a counseling session, a medical geneticist investigates individual's medical history and the family affected by the genetic disorder to order the right type of a genetic testing. Once a genetic diagnosis is achieved, in a post-counseling session, individual & the affected family will be taken through the implications of the result.

2 Carrier testing screens individuals for genetic mutations to reduce the risk that they'll pass down a genetic disorder to their children. If positive, couples will be given solutions to lower the chance of delivering a child with a genetic disorder. Carrier screening comes with two counseling sessions: Pre- and post-test counseling.

3 Clinicians submit individuals genetic raw data for previously undiagnosed cases. Data will be re-analyzed against GeneOriginS™ proprietary database with the goal of uncovering genes in complex cases. As an outcome, clinicians will be having a better understanding of the disease mechanisms which helps to Inform better clinical decisions.

4 Collaborate with GeneOriginS™ International scholars in rare disease research program. Talk to our experts, express your research interest, and join GeneOriginS™ community of gene discoverers.

How It Works



1

Talk to an Expert



2

Get Your Customized Service Solution



3

Save & Improve More Lives

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