The My Retina Tracker® Program provides open access, comprehensive, no-cost genetic testing and counseling to individuals who have a clinical diagnosis of an inherited retinal disease (IRD) and who live in the United States.

**HOW CAN I PARTICIPATE?**

This program is sponsored by the Foundation Fighting Blindness, and genetic counseling is provided by InformedDNA, a telephone-based counseling service.

The program provides patients with a 285 gene panel targeting relevant genes associated with an IRD. Unique features of the panel include full RPGR coverage, high resolution copy number variant detection and comprehensive coverage of IRD related non-coding variants.

A person with an IRD should speak to their retinal healthcare provider (ophthalmologist, retinal specialist, optometrist) and ask if they qualify for the test, and if so, ask their provider to order the My Retina Tracker Panel from Blueprint Genetics.

The health professional can find more information about the My Retina Tracker Program, and how to order the test, by visiting fightingblindness.org/open-access-genetic-testing-program.

If your doctor has questions, they should contact Blueprint Genetics directly.

Please do not try to order the test by contacting Blueprint Genetics yourself. Speak to your eye doctor as they are the only ones authorized to order the test.

**WHAT DOES THE TEST COST?**

There is no cost to the participant or their insurance company for the genetic test. Genetic counseling is also available at no cost through InformedDNA. You may need to cover the cost of a visit to speak with your clinician about the test, and for any tests they may want to do before ordering the test for you.

The My Retina Tracker Registry is a free service. Visit MyRetinaTracker.org.

For more information about the registry or genetic testing, contact Coordinator@MyRetinaTracker.org or 1-800-683-5555.

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WHO IS ELIGIBLE

FOR TESTING?

Any person with a clinical diagnosis of an IRD. The list includes retinitis pigmentosa, Leber congenital amaurosis, Stargardt disease, Usher syndrome, Best disease, choroideremia, achromatopsia and any of the other 25 covered diseases.

WHAT ARE THE BENEFITS?

With the increasing number of gene-specific therapies being developed, an accurate genetic diagnosis is very important for any person with an IRD.

This genetic test is specifically targeted to those with an IRD and is the most accurate, scientifically advanced, and highest quality test widely available to patients today.

Although it is not required for participation, this program offers an easy opportunity to join the My Retina Tracker Registry.

MY RETINA TRACKER REGISTRY

My Retina Tracker® Registry is a research database of people and families affected with an IRD. The purpose of the My Retina Tracker Registry is to accelerate the delivery of therapies for IRDs by determining the causes and prevalence of the different IRDs, supporting the research community to actively collaborate and promote research and development in the field of IRDs and efficiently enable people to connect to relevant clinical trials.

My Retina Tracker Registry is provided by the Foundation Fighting Blindness, a U.S.-based nonprofit organization with the mission to help find treatments and cures for all inherited retinal diseases.

WHY JOIN?

- The registry data drives more research.
- More registrants with a specific IRD can draw researchers’ focus and help support trial planning and funding.
- Registrants enter and control their own data and can withdraw at any time.
- Registrants have more access to announcements regarding focus groups, disease progression studies, genetic studies and clinical trials recruitment and updates.
- Your personal data and privacy are protected and only de-identified information is shared with researchers and clinicians.