WHAT IS THIS TEST?

myGenome with Diagnostic Interpretation is a genetic test that utilizes whole genome sequencing (WGS) with diagnostic interpretation to help identify a condition suspected to have a genetic cause. Additionally, the test includes information on Drug Sensitivities (Pharmacogenomics) and, for adults, a full myGenome report.

WHO IS IT FOR?

myGenome with Diagnostic Interpretation is for patients with a complex medical history or symptoms suggestive of a genetic condition, even if other genetic tests have shown a negative result. Patients may learn about genetic variants that explain signs and symptoms of their condition. Results can have implications for treatment decisions and outcomes, medical management options, as well as impact other family members. This product is available for adults (ages 18y+) and children (with parental consent).

WHY VERITAS?

Veritas is The Genome Company. Our mission is to deliver on the promise of the human genome, empowering people to maximize quality and length of life for themselves and their families.

As the first company to introduce whole genome sequencing and interpretation to consumers and their physicians, we lead by relentlessly pushing the boundaries of science and technology while driving down the cost of the genome.

Founded in 2014 by leaders in genomics from Harvard Medical School, we operate globally from offices in the U.S., Europe and China. We have been recognized twice by MIT Technology Review as one of the 50 Smartest Companies in 2016 and 2017, by Fast Company as one of the world’s most innovative health companies in 2018, and by CNBC as one of the Disruptor 50 Companies in 2018.
Pricing

It is highly-recommended that biological parental samples be tested when available as they can be helpful to interpret the patient’s genomic data. Please note: parental sample analysis is included in the test price, but a separate parental report is not provided.

New Patient: $2,999

Existing myGenome Customer: $2,400

Additional family member samples (blood relatives): Please inquire for additional cost

Technical Information

» Sequencing performed on the Illumina HiSeq X10 and NovaSeq 6000 systems

» Veritas follows Laboratory Developed Test (LDT) regulations performed in a CLIA-certified and CAP-accredited laboratory (CLIA ID: 22D2089381, CAP ID: 9289026)

» American College of Medical Genetics and Genomics (ACMG) guidelines are used for variant classification

» Pathogenic and Likely Pathogenic variants are confirmed with Sanger sequencing for the diagnostic portion of the test

» Large deletions and duplications, repeat expansions, certain inversions and deletions, translocations, and large structural rearrangements are not detected

» Only inherited (germline) variants are detected, and not somatic variants, mosaicism, or heteroplasmy

Learn More

If you have questions please contact us at 1 (888) 507-6619 or support@veritasgenetics.com