

UNCOVERING RARE OBESITY®

THE ONLY NO-CHARGE, COMPREHENSIVE GENETIC TESTING PROGRAM FOR RARE GENETIC DISEASES OF OBESITY

Genetic testing can be an essential tool in obesity management

All obesity is not the same. Certain forms of obesity are caused by genetic variants. Genetic testing can help you and your patients understand a primary cause of their obesity by identifying relevant genetic variants.

This knowledge can inform:

- Possible diagnosis
- · Appropriate disease management options
- Potential eligibility for clinical studies

Clinical guidelines for the assessment, treatment, and prevention of obesity recommend genetic testing to inform diagnosis and appropriate interventions in patients with early-onset, severe obesity and hyperphagia.^{1,2}

Individuals with these hallmark symptoms may be eligible for a no-charge genetic test. Rhythm Pharmaceuticals covers the cost of the test and provides sample collection kits. Patients are responsible for office visit, sample collection, and other costs.

Eligibility

To be eligible for testing through the Uncovering Rare Obesity program, patients must be located in the United States and its territories, AND:



ABOUT THE UNCOVERING RARE OBESITY PROGRAM



UncoveringRareObesity.com

your new destination to order kits, view results, and access educational materials.

References: 1. Cuda S, Censani M, O'Hara V, Browne N, Paisley J. Pediatric Obesity Algorithm eBook, presented by the Obesity Medicine Association. www.obesitymedicine.org/ childhood-obesity. 2020-2022. www.obesitymedicine.org/childhood-obesity (Accessed June 8, 2021). 2. Styne DM, Arslanian SA, Connor EL, et al. Pediatric obesity – assessment, treatment, and prevention: An Endocrine Society clinical practice guideline. J Clin Endocrinol Metab. 2017;102(3):709-757.

Program details



Testing conducted by a laboratory partner

DNA testing is conducted by

Extensive panel offers

clinical laboratory.

broad insights

PreventionGenetics, a CLIA-accredited

The test panel has been expanded from

40 genes to 79 genes and 1 chromosome

region, reflective of nearly all of the most

frequently tested genes associated with

obesity. Patients previously tested may

benefit from retesting, as it could offer

a test for Prader-Willi syndrome.

novel insight into their condition. The full

list of genes and region tested is available

on UncoveringRareObesity.com. This is not

Multiple sample collection options

Collection kits are available for use in the

office or to be sent to the patient's home.

blood or OCD-100 buccal swabs. If preferred,

OCD-100 buccal at-home kits are available to

send directly to patients. Spanish-language materials are also available upon request.

Samples may be collected in-office via



Easy access to test results

Test results will be available online and sent to you approximately 3 weeks after the lab receives the sample and completed forms.



Tailored support for results interpretation

The program provides you with access to a geneticist through our partner, PreventionGenetics, to help interpret results. For your patients, 2 sessions with licensed genetic counselors are available at no charge through our partner, PWNHealth, to provide guidance on the potential impact of genetic testing and to answer questions regarding results. Interpretation is available for multiple languages.

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Dedicated resources

Resources are available to support your ongoing needs following testing, including materials to help determine possible interventions. Disease education materials are also available for your patients.



UncoveringRareObesity.com

ORDER KITS / GET RESULTS / ACCESS RESOURCES

Contacts

FOR HEALTHCARE PROVIDERS

PreventionGenetics | 1-844-513-3994 Monday – Friday, 9am – 8 pm ET

FOR PATIENTS

PWNHealth | 1-888-494-7333 Monday – Friday, 9am – 5pm ET

GENERAL

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