Cure HHT is collaborating with the Rare Genomes Project, a research program that performs genetic sequencing on families with suspected genetic diseases.

**ABOUT THE PROJECT**

The Rare Genomes Project is a patient-driven research study led by genomics experts and clinicians who believe that the latest advances in genomic sequencing are changing medicine and that they should be accessible to families with rare and undiagnosed conditions. The aim of the project is to use genomic sequencing to look for the genetic cause of rare diseases in families.

**WHAT IS INVOLVED IN THIS STUDY?**

- Participation is remote and free of cost. Eligible families will be asked to provide relevant medical information and to donate a blood sample using a collection kit mailed to their home.
- If a genetic variant that may be the cause of HHT is found, the Rare Genomes Project will contact you and work with your doctor to confirm results.

This research process takes longer than routine genetic testing and not all families will have a genetic change identified.

**WHO CAN PARTICIPATE?**

The Rare Genomes Project is looking to enroll individuals from families where a genetic cause has not been identified and who either:
- Cannot get routine genetic testing or
- Have had negative or inconclusive prior genetic testing results

Participants must also:
- Live in the United States
- Be English- or Spanish- speaking

**HOW TO APPLY**

Visit raregenomes.org/how-it-works or call 617-714-7395 to learn more and apply. Only one member of the family should submit an application.

Questions? Email us at researchnetwork@curehht.org