National Consumer Center for Genetics Resources & Services

Empowering the advocacy community through education and policy

In collaboration with disease-specific advocacy organizations and family groups, the National Consumer Center for Genetics Resources & Services (NCCGRS) develops and makes accessible informational materials and educational programs for consumers. Further, the Center will bring consumers and their concerns to the attention of policymakers and government agencies. The National Consumer Center for Genetics Resources and Services seeks to promote the collaboration required to revolutionize networks and systems to serve the public’s needs.

The NCCGRS includes the following programs:

• **Access to Credible Genetics Resources Network:** Establishing standards for accurate and credible information on genetic conditions and providing tools for the development of quality materials
• **Community Centered Family Health History:** Engaging diverse communities in the creation of family health history resources
• **Consumer-Focused Newborn Screening Projects:** Generating models to maximize benefits in newborn screening

The Center focuses on ensuring the consumer voice is heard and on improving services for all who need them. Every person will need genetics resources and services some day. The Center will work proactively to focus the future of the field on the issues that matter most to the consumer and make sure these services are accessible, affordable, and of high quality.

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