NEWLY DIAGNOSED?

We are here for you and want you to know that you are not alone. Welcome to the FamilieSCN2A Foundation!

Who Are We?

"Families" is part of our name for a reason. Rare and devastating, SCN2A-related disorders affect the entire family. Our team of leaders strive every day and in every way to improve the lives of not only the patients, but the entire family.

Our VISION is a world with effective treatments and cures for all SCN2A-related disorders.

Our MISSION is to accelerate research, build community and advocate to improve the lives of those affected by SCN2A-related disorders around the world.

PROGRAMS AVAILABLE

Global SCN2A Support Network is a private network of Facebook pages established by the FamilieSCN2A Foundation for parents, caregivers, patients and families of children with SCN2A related disorders.

SCN2A Family Event Grants serve to intentionally bring SCN2A families around the world together to cultivate a stronger community and deepen relationships.

SCN2A Patient Assistance Grants are in place to offer small grants to patients with SCN2A Disorders for necessary medical equipment, therapy devices, and educational aids associated with these conditions that are not covered through private insurance or other assistance programs.

Birthday Club is a way for us to show our love and celebrate our kids on their special day!

Newsletter and Emails help you to stay current on all that the Foundation is up to, SCN2A related research, and important SCN2A news by signing up to receive our newsletter and emails.

Warrior Wednesdays we welcome you to post pictures of your Warrior and share a story about how they / you are “fighting on”.

We offer several programs to our global community. Visit our website to learn more: www scn2a.org