The FamilieSCN2A Foundation is dedicated to finding effective treatments and a cure for SCN2A disorders. We will accomplish this while improving the lives of those affected by SCN2A disorders through clinical research, effective treatments, public awareness, early detection, patient advocacy, and family support. We can’t do this alone and have built a large community of fierce families to fight with. Each of you play a vital role in pushing the needle closer to a cure.

The SCN2A Virtual Conference is a two part series hosted by the FamilieSCN2A Foundation and the Simons VIP Connect. The focus is on educating families, physicians, and other community members on the advancements made in the field. We thank Simons VIP Connect for their
partnership and drive on all genetic research including SCN2A.

Part One will be held on October 6, 2018 will spotlight the latest exciting discoveries in basic research, drug development and improvements in clinical treatments. Our presenters include Alfred L. George, Jr., MD; Keith A. Coffman, MD; Stephan J. Sanders, BMBS (MD) PhD; Wendy K. Chung, MD, PhD; and Heather E. Olson, MD. These brilliant scientists are just as passionate about SCN2A as we are!

**DATE:** Saturday, October 6, 2018  
**TIME:** 12:00 p.m. - 2:30 p.m. (Eastern Daylight Time)  
**LOCATION:** wherever you are!

Register Here

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Save the Date: November 3, 2018  
Part Two will spotlight therapies and alternative medical approaches.  
Details coming soon!