

WELCOME to All Things Kabuki and the Kabuki Syndrome family! We're glad you're here and look forward to getting to know you. Whether your child was recently diagnosed, or you've just found our community, you will find this to be a place of encouragement, love and support.

ALL THINGS KABUKI

AWARENESS. EDUCATION & SUPPORT FOR KABUKI SYNDROME

All Things Kabuki is a volunteer run 501(c)(3) non-profit corporation based out of Wasilla, Alaska. ATK was founded in 2013 and is the only U.S. based patient advocacy group supporting individuals and families affected by Kabuki Syndrome around the globe. We have a wonderful team, all caregivers and relatives of children with Kabuki, ready and willing to help. You can learn more about our volunteers on our website.

Enclosed in this *Welcome Folder* are several resources we hope you'll find helpful, some of which include: our programs, parent tips, information on research and survey opportunities, and awareness resources! ATK offers a variety of different resources to the community, all of which can be ordered at no cost via our website or by email. We encourage you to use these tools to join our efforts to raise awareness of Kabuki Syndrome.

As a volunteer run organization, we rely on the community to partner with us to not only promote awareness, but to raise funds to support the programs we offer and to provide advocacy and awareness resources at no cost to the global community. There are many ways friends and family can contribute to our mission, whether it be through Facebook fundraisers, employer matching donations, local fundraising efforts, or direct contributions through our website. All donations made to ATK are tax deductible. You can find additional ways to support ATK on our website.

If you are a parent or primary caretaker of an individual with Kabuki Syndrome, we encourage you to join ATK's *Kabuki Syndrome Parents* Facebook group. This is a *parent-only* forum established to offer a private, safe place for parents to talk, laugh and vent. Don't forget to check out and follow our Facebook page **All Things Kabuki** for frequent updates and be sure to sign up for our electronic newsletter.

If we can do anything to help your family on your Kabuki journey, please don't hesitate to contact us.

Best wishes,

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Rene King Founder & CEO

OUR MISSION IS TO RAISE AWARENESS, INCITE RESEARCH AND SUPPORT INDIVIDUALS AND FAMILIES AFFECTED BY KABUKI SYNDROME.