

ONLINE EVENT

MARCH 27, 2024
9:30am-12pm



A DISCUSSION ABOUT FAMILY EXPERIENCES AFTER NEWBORN SCREENING

FOR WHO?

Parents or caregivers of children who had a medical condition diagnosed from newborn screening (bloodspot, hearing, or cardiac tests) in DE, DC, MD, NJ, NY, PA, PR, USVI, VA, or WV

WHAT?

A 2.5 hour online conversation with a small group of other families

WHY?

To learn from families about what aspects of their family's life were impacted by newborn screening, how families are doing years after the diagnosis, and how they can be better supported

Spanish
interpretation
and closed
captioning

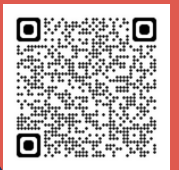
If you are interested, please fill out the form and we will contact you if you are selected to participate (click below or use QR code)

<https://forms.gle/2zFyPPmQztA5Wzgb6>



Families who are selected for the session will receive a stipend for their time

¿Interesado?
Haga clic
aquí para
español



For More Information about Newborn Screening:

<https://www.babysfirsttest.org/newborn-screening/about-newborn-screening>

 NYMAC@ferre.org

 [NYMACgenetics.com](https://www.nymacgenetics.com)

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