



NORD®
National Organization
for Rare Disorders

EMPOWER CHANGE

Your Voice Matters in Building
a Rare Disease Advisory Council



The National Organization
for Rare Disorders (NORD®)
is working to establish
Rare Disease Advisory
Councils (RDAC) in every
state through Project RDAC.

**Currently, there are
RDACs in 28 states.**



What is an RDAC?

An RDAC serves as a state-level advisory body, providing a voice for the rare disease community in state government. Despite the diverse nature of rare diseases, this community frequently encounters common challenges such as accessing specialists, medications, facing misdiagnoses, and finding resources. RDACs offer a platform for rare disease patients to directly engage with lawmakers, fostering opportunities for positive legislative changes.



Discover Your Role Today!

Join the coalition of rare disease patients, experts, and advocates like you to push lawmakers to create a Rare Disease Advisory Council.

- Meet other rare disease community members in your state
- Discuss the unique needs and challenges that the rare disease community faces
- Become an active participant in rare disease policy advocacy efforts before your state lawmakers



For more information on
Project RDAC, please contact
RDAC@rarediseases.org or
visit **rarediseases.org**