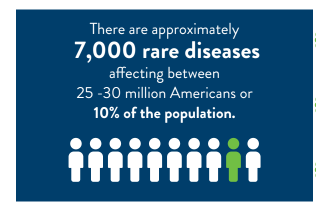
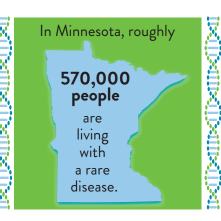
## Rare isn't so rare.

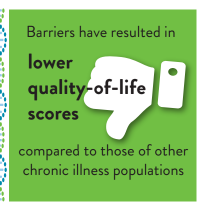


Any disease, disorder, illness, or condition affecting fewer than 200,000 people in the U.S. is considered **rare**.

Minnesota Rare Disease Advisory Council is a result of the grassroots efforts of Minnesota patients and families whose lives have been affected by rare disease. The Council was established in 2019 by the Minnesota state legislature to ensure that every patient diagnosed with a rare disease has access to a timely diagnosis, appropriate care, and an effective treatment. The Council consists of rare disease experts from across health systems and aligned disciplines who work tirelessly to create a strong responsive, rare disease community.







Average time to diagnosis for a rare disease is **7-8 years.** 



Individuals are misdiagnosed **2-3 times** 



Individuals eventually diagnosed with a rare disease see an average of **eight clinicians** prior to diagnosis



Only 5%
of rare disease patient
populations have an
FDA approved treatment



Rare disease
diagnoses are diverse;
however, patients'

barriers to care
are common.

Research footnotes can be found at www.cbacraredisease.org



Envisioning a world where every Minnesota citizen living with a rare disease has access to a timely diagnosis, comprehensive care, and an effective treatment.

