




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SODRE FOUNDATION

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Hosted by SODRE Foundation



LOS VERDES GOLF COURSE
JULY 24, 2024



Supporting
Opportunities
Dreams
Rare disease community
Education + advocacy

ABOUT US

SODRE Foundation was founded by a junior golfer named Audrey Chang. Audrey's journey began with a personal story of love and determination.

Witnessing her older sister, Sophia's struggles with DDX3X syndrome, a rare genetic disorder, Audrey decided to take action knowing that treatment and therapy options could improve the quality of her sister's life. She combined her love and passion for golf with her commitment to make a difference, leading to the establishment of SODRE Foundation in 2023.



MISSION & GOALS

Our mission at SODRE Foundation is **S**upporting **O**pportunities and fulfilling **D**reams for the **R**are disease community through **E**ducation and advocacy.

At Sodre Foundation, our goal is to raise awareness, educate, and raise money for various rare diseases through organized events and fundraisers.

ABOUT DDX3X

DDX3X Syndrome is a rare disease caused by a spontaneous mutation within the DDX3X gene at conception or can be inherited. The syndrome was discovered in the United States in 2014 and primarily affects girls due to its location on the X-chromosome, though there are some boys it has affected as well. Although it has only been identified in about 1000 individuals, doctors believe it is the cause of 1 to 3 percent of all intellectual disabilities in females.

Not all individuals with DDX3X Syndrome are affected in the same ways, however, the following are common symptoms:

- Intellectual disability
- Developmental delays
- Low muscle tone/hypotonia
- Difficulty with speech
- Epilepsy/seizures
- Movement disorders
- Abnormalities of the brain
- Microcephaly
- Scoliosis

All proceeds from this event will be donated to the DDX3X Foundation. The ultimate goal of the DDX3X Foundation is to raise money to fund research to accelerate brain function in individuals affected by DDX3X Syndrome through advances in cell and gene therapy and pharmaceuticals.