Contact Us

For additional support and information, contact the CHARGE Syndrome Foundation at info@chargesyndrome.org

A better world for people with CHARGE syndrome.

The CHARGE Syndrome Foundation champions the lifelong potential of people with CHARGE syndrome through outreach, education and research.
What Is CHARGE Syndrome?

CHARGE is a complex genetic syndrome that includes a range of physical and health challenges that vary from child to child. It is the most common genetic cause of congenital deaf-blindness, with an incidence of about 1 in 12,000 births. In most cases, there is no family history of the disorder or similar conditions.

Birth defects and medical features may include:
- Coloboma of the eye
- Choanal atresia or stenosis
- Facial palsy
- Tracheal or esophageal abnormalities
- Swallowing difficulties
- Breathing issues
- Ear malformations (outer and inner ear)
- Heart defects
- Additional organ system problems

CHARGE Syndrome Also Can Affect Sensory Systems Including:
- Vision
- Hearing
- Balance
- Behaviors
- Sleep
- Feeding and digestion

Every individual with CHARGE syndrome is unique in their needs and abilities. There is no predetermined limit to an individual’s capacity for learning and educational achievement.

The mission of the CHARGE Syndrome Foundation (CSF) is to create a better world for people with CHARGE.

The CSF provides support and outreach to individuals with CHARGE, their families, and professionals. The CSF gathers, develops, maintains, and distributes information about CHARGE, and promotes awareness and research.

Local, national, and international partners provide an array of services to support the CHARGE syndrome community. We look forward to supporting your family every step of the way.