

Join the Conversation

chromodisorder.org, Twitter, Facebook, YouTube and our other social media sites offer individuals and families coping with a rare disorder diagnosis a chance to reach out 24 hours a day.



Chromosome Disorder Outreach, Inc.

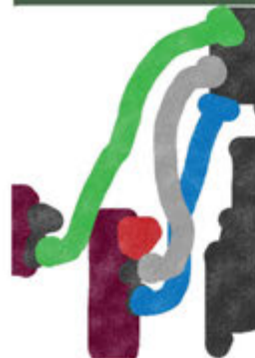
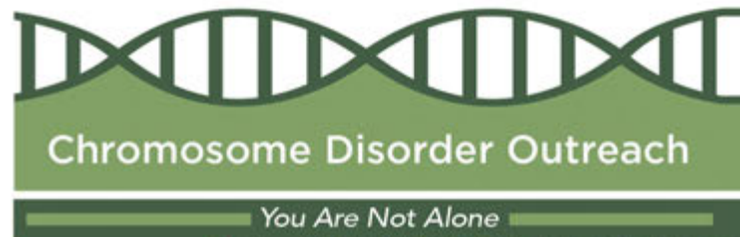
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Family Helpline
561.395.4252

info@chromodisorder.org
www.chromodisorder.org

*Affiliated with Genetic Alliance,
NORD, Rare Disease Day.*

*CDO highly respects each member's personal data.
Members direct their own privacy settings and
confidentiality is never compromised.*



*Since 1992, CDO has been
working to improve the lives of
children and adults coping with
any rare chromosome disorder.*

Help Raise Awareness

Visit

chromodisorder.org

and learn more about how you can spread
the word to raise awareness.

Chromosome Disorder Outreach, Inc. (CDO)
is a non-profit 501 C3 organization.

100% of contributions are used to support
individuals and families coping with
a rare chromosome disorder diagnosis.

CDO does not employ professional fundraisers.
Donations are fully tax deductible as provided by law.
Florida Registration #CH11200

Cover Art: Kyle, age 5, Gus, age 9, Mark, age 7





BRAVE

YOU ARE NOT ALONE



"I can still remember how lonely and afraid I was when Steven was diagnosed. CDO has removed that loneliness for countless others. Thank you. No other words but thank you."

**– Joan Rhoades
CDO founding member**

Chromosome Disorder Outreach Inc. is a non-profit organization founded in 1992 by a group of seven parents raising children born with rare chromosome deletions. Today our members number in the thousands and are diagnosed with a wide range of chromosome disorders including deletions, duplications, trisomies, inversions, translocations, and rings. Sometimes these disorders are so unusual that doctors tell us, "You're the only one out there."

But with CDO, you are not alone. Our mission is to provide support and information to families and individuals diagnosed with chromosome disorders.

CDO's personalized networking programs can help those newly diagnosed locate others in their area and around the world who share common challenges and successes. Reach out through our social media applications and connect whenever you choose – parent to parent support is available online 24 hours a day. CDO keeps members updated on new research opportunities as well as the latest in rare chromosome publications. Our library contains over 4,000 articles. Newsletters and an email database allow us to share research opportunities and other information relative to the community.

CDO encourages awareness and education of the general public. In June 2014, CDO helped support the first annual Rare Chromosome Disorder Awareness Week. This grassroots effort will grow year after year and enable more families and individuals to learn more about CDO and rare chromosome disorders.

Become a Member Today!

Membership is free and comes with these benefits:

- Personalized networking programs
- Database of registered disorders
- Closed Facebook group
- Quarterly online newsletters
- List of support groups for particular disorders
- Library of online scientific information
- "Ask the Doctor" archive and medical query resource



COMPASSIONATE