Miracle Flights (MiracleFlights.org)
Provides financial assistance to low-income children for commercial air travel to obtain special medical care.

The Sing Me a Story Foundation (SingMeAStory.org)
Combines the imaginations of children with the talents of songwriters to create stories and songs that bring joy to all those involved.

Good Days (GoodDays.org)
Works to improve the health and quality of life of patients with chronic disease, cancer, or other life-altering conditions. The organization may help provide families with financial and travel assistance.

The HealthWell Foundation (HealthWellFoundation.org)
Independent non-profit dedicated to reducing financial barriers to care for underinsured Americans with chronic and life-altering medical conditions. HealthWell offers a financial lifeline to adults and children who desperately need critical medical treatments but can’t afford them by assisting with their cost-sharing obligations.

National Organization for Rare Disorders (RareDiseases.org)
Provides many resources for people living with rare diseases, their families and other advocates, including but not limited to:

- Co-pay Program*: assists eligible individuals who have health insurance to cover health insurance deductibles, copayments and coinsurance costs associated with the care of chronic granulomatous disease.
- Medical Assistance Program*: assists eligible individuals who are uninsured or for whom coverage has been denied, with out-of-pocket costs for medical expenses such as medical visits, laboratory and diagnostic testing, physician prescribed medical foods & supplements, travel to and from a disease-related medical appointment and other expenses for the care of chronic granulomatous disease.

Global Genes (GlobalGenes.org)
Develops educational resources, programs, and events that unite patients, advocates, and industry experts.

- Global Genes RARE Patient Advocacy Summit: The largest rare patient advocate event worldwide.
- Scholarship to the RARE Patient Advocacy Summit: With the help of generous sponsors, advocates from all over the globe are afforded travel and lodging scholarships to attend the Summit. E-mail events@globalgenes.org for scholarship information.

About Horizon
Horizon is focused on researching, developing and commercializing medicines that address critical needs for people impacted by rare and rheumatic diseases. Our pipeline is purposeful: we apply scientific expertise and courage to bring clinically meaningful therapies to patients. We believe science and compassion must work together to transform lives. For more information on how we go to incredible lengths to impact lives, please visit www.horizontherapeutics.com, follow us @HorizonNews on Twitter, like us on Facebook or explore career opportunities on LinkedIn.

Connection to Chronic Granulomatous Disease and Rare Disease Support and Services
Horizon provides support to many organizations that provide a wide range of crucial programs and services for people living with chronic granulomatous disease and other rare diseases.

For information about each of the above programs, send an e-mail to CGD@rarediseases.org.

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Support and Assistance

Patient Support and assistance in place so that, where eligible, patients are able to receive our medicines regardless of their ability to pay.*

Connection online and in person with other families living with CGD to help offer tips from their experience, answer questions and support through life’s transitions

CGD Nurse Educators available to those who are prescribed our medicine at every step of their journey. These CGD specialists provide personalized support and individual guidance about CGD and injection training

* Assumes utilization of Horizon financial support programs

Research and Development

FURTHERING EXPERTISE AND DIAGNOSIS

Sponsor

the creation of the Jeffrey Modell Foundation Diagnostic and Research Centers for Primary Immunodeficiencies at St. Louis Children’s Hospital and at the University of Virginia to ensure cutting-edge primary immune deficiency diagnostics are available to more children

Support

the development of a Journal of Pediatric Infectious Disease Society supplement on understanding CGD and the Clinical Immunology Society’s development of a CGD specific educational video series based on the supplement

ADVANCING NEW RARE DISEASE OPTIONS

$225 million

in research and development expenses incurred by Horizon in 2017 for rare and rheumatic diseases**

Multiple clinical programs:

exploring new compounds

exploring new uses for our current medicines

** Includes costs associated with the acquisition of River Vision Development Corp.

SUPPORTING EXTERNAL RESEARCH

$3 million

dollar gift to Children’s National Rare Disease Institute to establish an expert care team in rare disease

$10 million

in funding for 29 external studies evaluating our medicine for CGD to help address a range of potential diseases

$5.5 million

dollar donation of our medicine for clinical research at the National Institutes of Health (NIH)

Community Engagement

CGD Connections™ website and Facebook page co-created with people living with CGD and their families provides education and resources

• Educational fact sheets and animations
• Expert videos
• Career and school resources
• Caregiver support

annual events

supported and attended to help bring people together

• Jeffrey Modell Foundation Kids Days
• Immune Deficiency Foundation Walks
• Special CGD-specific sessions at Immune Deficiency Foundation retreats and National Conferences

LivingWithCGD.org

a resource made possible through an unrestricted grant to the Immune Deficiency Foundation featuring videos and blog posts from CGD community members sharing their tips, experience and the latest news

20+

rare disease events sponsored and attended each year by Horizon employees

50+

patient advocacy group partnerships globally, including leading primary immunodeficiency advocacy organizations

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