



The Global Genes RARE Foundation Alliance is made up of over 750 disease foundations that have committed to collaborating with Global Genes and other nonprofit foundations in order to create a stronger, collective voice in the rare disease community.

[#Bold Lips For Sickle Cell](#) – Sickle Cell Disease

[11q Research & Resource Group](#) – Jacobsen Syndrome, 11q Chromosome

[1p36 Deletion Support & Awareness](#) – 1p36 Deletion Syndrome

[22q 11 Ireland support group](#) – 22q11.2 deletion syndrome

[4p- Support Group](#) – Wolf-Hirschhorn Syndrome and related 4p conditions

[5p-Society](#)– 5p- Syndrome, Cat Cry Syndrome, Cri du Chat Syndrome

[17q12 Foundation](#) - 17q12 Deletions and Duplications

[A Breath Of Hope Foundation For NMO](#) - Neuromyelitis Optica

[A Foundation Building Strength for Nemaline Myopathy – Nemaline Myopathy](#)

[A Nonprofit Group Enriching Lives \(ANGEL AID\)](#) - Multiple rare diseases

[Aaron's Ohtahara](#) – Ohtahara Syndrome

[Acid Maltase Deficiency Association](#)– Acid Maltase Deficiency, Pompe's Disease

[Acromegaly Community](#) – Acromegaly and Gigantism

[Acromegaly Ottawa Awareness & Support Network](#) - Acromegaly

[Acoustic Neuroma Association](#) – Acoustic Neuroma

[ADCY5.org](#) – ADCY5 Mutation

[Addi & Cassi Fund](#) – Niemann Pick Type C

[ADNPkids](#) – ADNP Syndrome, Helsmoortal\_Van Der AA Syndrome

[Adrenal Alternatives Foundation](#) - Adrenal Diseases

[Adrenal Insufficiency United](#) – Adrenal Insufficiency

[Adult Polyglucosan Body Disease Research Foundation](#) (APBDRF) – APBD

[Advancing Sickle Cell Advocacy Project, Inc.](#) – Sickle Cell Disease

[Advocacy & Awareness for Immune Disorders Association](#) – Primary Immunodeficiency Diseases (PID), Hereditary Angioedema, Pediatric Acute onset Neuropsychiatric Syndrome (PANS), Autoimmune diseases

[AGO2 Association](#) - Lessel-Kreienkamp Syndrome (AGO2 gene)

[AHC Federation of Europe](#) – Alternating Hemiplegia of Childhood

[Aicardi-Goutieres Syndrome Association of the Americas](#) (AGSAA)- Aicardi-Goutieres Syndrome

[Aidan Jack Seeger Foundation](#) – Adrenoleukodystrophy

[AKU Society](#) – Alkaptonuria

[AKU Society North America](#) – Alkaptonuria

[Alagille Syndrome Alliance](#) – Alagille Syndrome

[ALD Alliance](#) - ALD Adrenoleukodystrophy

[Alect2 Alliance](#) – Amyloidosis Alect2

[ALD Connect](#) – Adrenoleukodystrophy

[All Things Kabuki](#) – Kabuki Syndrome

[Alliance for Cryoglobulinemia](#) – Cryoglobulinemia

[Alliance for Safety Awareness for Patients](#) – Hospital-Acquired Necrotizing Fasciitis

[Alpha-1 Foundation](#) – Alpha-1 Antitrypsin Deficiency

[Alport Syndrome Foundation](#) – Alport Syndrome

[ALS Ride For Life](#) - ALS

[Alström Angels](#) – Alström Syndrome

[Alström Syndrome International](#) – Alström Syndrome

[Alternating Hemiplegia of Childhood Foundation](#) – Alternating Hemiplegia of Childhood

[AMENSupport](#) – Multiple Endocrine Neoplasia

[American Behcet’s Disease Association](#) – Behcet’s Disease

[American Brain Tumor Association](#) - Brain Tumors

[American Partnership for Eosinophilic Disorders](#) – Eosinophilic Disorders

[American Society of Cell and Gene Therapy](#)

[American Porphyria Foundation](#) – Porphyria Disease

[Amour Fund of Alpha Epsilon Omega Foundation](#) – Congenital Disorders of Glycosylation (CDG)

[Amyloidosis Foundation, Inc.](#) – Amyloidosis

[Amyloidosis Research Consortium](#) – Amyloidosis

[Amyloidosis Support Groups Inc.](#) - Amyloidosis Support Groups throughout the United States

[AnCan\(formerly Answer Cancer Foundation\)](#) - Sarcoidosis, Renal Medullary Cancer, Male Breast Cancer

[Angelman Biomarkers and Outcome Measures Alliance](#) (A-BOM)

[Angelman Syndrome Foundation](#) - Angelman Syndrome

[Angels with Missing Pieces](#) – 22Q Deletion Syndrome

[Angioma Alliance](#) – Cavernous Angioma

[Appendix Cancer / Pseudomyxoma Peritonei Research Foundation \(ACPMP\)](#)– Pseudomyxoma Peritonei

[Aplastic Anemia & Myelodysplasia Association of Canada \(AAMAC\)](#) - Aplastic anemia, myelodysplastic syndrome and paroxysmal nocturnal hemoglobinuria

[Aplastic Anemia and MDS International Foundation](#) – Aplastic anemia, Myelodysplastic syndromes, Paroxysmal nocturnal hemoglobinuria

[APFED - American Partnership for Eosinophilic Disorders](#) - Eosinophilic related disorders

[Arachnoiditis Society for Awareness and Prevention](#) – Arachnoiditis

[Arachnoiditis and Chronic Meningitis Collaborative Research Network](#)

[ARPKD/CHF Alliance](#) - Autosomal Recessive Polycystic Kidney Disease and Congenital Hepatic Fibrosis

[Asociación Española De Amiloidosis](#) - Amyloidosis

[Asociación Puertorriqueña de Hemofilia y Condiciones de Sangrado \(APH\)](#) - Hemophilia

[Association for Creatine Deficiencies](#) - Creatine Transport Deficiency, Guanidinoacetate Methyltransferase Deficiency, L-Arginine: Glycine Amidinotransferase

[Association for Glycogen Storage Disease](#) - Glycogen Storage Disease

[Association of Peyronie's Disease Advocates](#) - Peyronie's disease

[AT Children's Project](#) - Ataxia-Telangiectasia

[Ataxia a Fi - Ataxia and Me](#) - Ataxia

[Atlanta S.C.A.R., Inc](#) - Sickle Cell Disease

[A Twist of Fate](#) - Arterial tortuosity syndrome

[Atypical HUS Foundation](#) - Atypical Hemolytic Uremic Syndrome

[Autoimmune Encephalitis Alliance](#) - Autoimmune Encephalitis

[Autoimmune Hepatitis Association](#) - Autoimmune Hepatitis

[Autoinflammatory Alliance](#) - Cryopyrin-Associated Periodic Syndromes, Neonatal-Onset Multisystem Inflammatory Disease, Chronic Infantile Neurological Cutaneous and Articular Syndrome, Familial Cold Autoinflammatory Syndrome

[Aux Pas Du Coeur](#) - GNE Myopathy, Duchenne Muscular Dystrophy

[Avery's Angels Gastroschisis Foundation](#) - Gastroschisis Disease

[Avery's Hope](#) - GI Diseases

[Aware of Angels](#) - Multiple Rare Diseases

[Axenfeld-Rieger Foundation](#) - Axenfeld-Rieger Syndrome

[AXSL Rare Research Endowment Foundation](#) - Bohring-Opitz, Shashi-Pena and Bainbridge Ropers Syndrome

[AXYS](#) - One or more extra X and/or Y chromosomes

[Bardet Biedl Australia](#) - Bardet-Biedl Syndrome

[Bardet Biedl Families](#) - Bardet Biedl Syndrome

[Barth Syndrome Foundation](#) - Barth Syndrome

[Batten Disease Support and Research Association](#) - Batten Disease

[Bcureful](#) - Tuberous Sclerosis Complex

[Beautiful You MRKH Foundation](#) - Mayer-Rokitansky-Küster-Hauser Syndrome

[Beck-Fahrner Syndrome Foundation](#) - Beck Fahrner Syndrome, TET3 Deficiency

[Beckwith-Wiedemann Children's Foundation International](#) - Beckwith-Wiedemann Syndrome

[Ben's Dream](#) - Sanfilippo Syndrome

[Ben's Friends](#) - All Rare Diseases

[Bethany's Hope Foundation](#)

[Beyond Batten Disease Foundation](#) - Batten Disease

[BlackSwan Foundation](#) - Multiple Rare Diseases, Switzerland

[Bladder Exstrophy Research Foundation](#) - Bladder Exstrophy

[Bleeding Disorders Alliance Illinois](#) - Inherited Bleeding Disorders  
[Blind Children's Learning Center](#) - Multiple Rare Diseases  
[Bloom Syndrome Association](#) - Bloom Syndrome  
[Bohring-Opitz Syndrome Foundation, Inc.](#) - Bohring-Opitz Syndrome  
[Boomer Esiason Foundation](#) - Cystic Fibrosis  
[Born a Hero](#) - Pfeiffer Syndrome  
[BPAN Warriors](#) - Beta-Propeller Protein-Associated Neurodegeneration  
[Brandon's Battle Foundation For SKS Kids](#) - Smith Kingsmore Syndrome

[Brave Minds Project](#) - Brain and Brainstem Conditions  
[Breathe Support Network](#) - Pulmonary Fibrosis  
[Bridge the Gap](#) - SYNGAP1  
[CACNA1A](#) - CACNA1A related diseases - epilepsy, ataxia, hemipelegic migraines, IDD  
[CADASIL Together We Have Hope](#) - Cerebral Autosomal Dominant Arteriopathy Subcortical Infarcts Leukoencephalopathy  
[California Action Link for Rare Diseases Inc. \(CAL RARE\)](#) - All Rare Diseases  
[Care Beyond Diagnosis](#) - Multiple Rare Diseases  
[Cailee's Corner](#) - Familia Exudative Vitreopathy (FEVR)  
[Campbell Burns Metabolic Trust](#) - Metabolic Disorders  
[Canadian Association of Pompe](#) - Pompe Disease  
[Canadian Fabry Association](#) - Fabry Disease  
[Canadian MPS Society](#) - Mucopolysaccharide and Related Diseases  
[Canadian Organization for Rare Disorders](#) - Multiple Rare Diseases  
[Canadian PKU & Allied Disorders](#) - Phenylketonuria  
[Canadian Pituitary Patient Network](#) - Pituitary and Adrenal disorders  
[Care4ASH1L](#) - ASH1L related disorders  
[CARES Foundation](#) - Congenital Adrenal Hyperplasia  
[Caring Voice Coalition](#) - Multiple Rare Diseases  
[Carion Fenn Foundation](#) - Syringomyelia, Arnold Chiari Malformation  
[Castleman Disease Collaborative Network](#) - Castleman Disease  
[Cauda Equina Foundation, Inc.](#) - Cauda Equina Syndrome and associated disorders  
[CCHS Family Network /CCHS Foundation](#) - Congenital Central Hypoventilation Syndrome  
[CDG Care](#) - Congenital disorders of glycosylation  
[CDH International](#) - Congenital Diaphragmatic Hernia  
[CDKL5 Canada](#)  
[CDKL5 Research Collaborative](#)  
[Center for Chronic Illness](#)  
[Central California Hemophilia Foundation](#) - Inherited Bleeding Disorders  
[CFC International](#) - Cardio-Facio-Cutaneous Syndrome  
[CHAMP1ResearchFoundation](#) - CHAMP1

## [Champions Action Network](#)

[Chasing Life Project](#) – Multiple Rare Diseases

[Charcot-Marie-Tooth Association](#) - Charcot-Marie-Tooth

[Chelsea's Hope Lafora Children Fund](#) - Lafora Disease

[Cheyanna's Champions 4 Children \(CC4C\)](#)– Multiple Rare Diseases & Undiagnosed

[chILD Foundation](#) – Interstitial Lung Disease

[Child Neurology Foundation](#) – Childhood Neurological Disorders

[Children's Alopecia Project](#) – Alopecia

[Children's Cardiomyopathy Foundation](#)– Pediatric Cardiomyopathy

Children's Cerebral Palsy Movement – Multiple Rare Diseases

[Children's Craniofacial Association](#) – Craniofacial Disorders

[Children's Gaucher Research Foundation](#) – Gaucher Disease Types 2/3

[Children's Tumor Foundation](#) – Neurofibromatosis Disease

[China-Dolls Center for Rare Disorders](#) - Rare Bone Disorders

[Chinese Organization for Rare Disorders](#) – Multiple Rare Diseases

[Chion Foundation](#)– Prader-Willi Syndrome

[Chive Charities](#) – Multiple Rare Diseases

[Cholangiocarcinoma Foundation](#)

[CHOPS Syndrome Global](#) - Chops Syndrome

[Chordoma Foundation](#) – Chordoma, Sarcoma

[Choroideremia Research Foundation](#) – Choroideremia

[Chronic Intractable Pain And You Sites, Inc.](#) – Chronic and/or Intractable Pain

[Chronic Pain Partners](#)

[Cincinnati Comprehensive Sickle Cell Center at Cincinnati Children's Hospital](#)– Sickle Cell Disease

[Citizens United for Research in Epilepsy](#) – Epilepsy

[CLOVES Syndrome Community](#) – CLOVES Syndrome

[CMCD Foundation](#) - Cortical Malformations and Cephalic Disorders

[CMTC-OVM US](#) – CMTC (Van Lohuizen Syndrome)

[Coalition Duchenne](#) – Duchenne Muscular Dystrophy

[Coalition to Cure CHD2](#) - Find a cure for CHD2-Related Epilepsy and Autism

[Coffin-Siris Syndrome Foundation](#) - Coffin-Siris Syndrome

[Cold Agglutinin Disease Foundation](#) - Cold Agglutinin Autoimmune Hemolytic Anemia-Acquired

[Colie Creations, Inc.](#) - Cystic Fibrosis, Transplant, Chronic Illness

[COMBINEDBrain](#) - General Support Organization

[Complex Child E-Magazine](#) – Multiple Pediatric Rare Diseases

[Congenital Hyperinsulinism International](#) – Congenital Hyperinsulinism

[Connecting Families Urea Cycle Disorders Foundation](#) - Urea Cycle Disorder

[Connor B. Judge Foundation](#) - NMO

[Cook for Love, Inc.](#) - PKU and other IEM that require a low protein diet

[Coriell Institute for Medical Research](#) – Multiple Rare Diseases

[Cornelia de Lange Syndrome Foundation](#) – Cornelia de Lange Syndrome

[Costello Syndrome Family Network](#) – Costello Syndrome

[Courageous Faces Foundation](#) - Multiple Rare Diseases

[CPRS Forum](#) – Complex Regional Pain Syndrome

[CRMO Foundation](#) - Chronic Recurrent Multifocal Osteomyelitis

[CSNK2A1 Foundation](#)– Okur-Chung Neurodevelopmental Syndrome (OCNDS)

[CTNNB1 Syndrome Awareness Worldwide](#) - CTNNB1 Syndrome

[Cure AHC](#) - Alternating Hemiplegia of Childhood

[CureARS](#) - Mitochondrial Aminoacyl-tRND Synthetase Research & Awareness

[Cure CADASIL](#) -CADASIL

[Cure CMD](#) – Congenital Muscular Dystrophy

[Cure CMT4J/ Talia Duff Foundation](#)– Charcot Marie Tooth Disease, Type 4J (CMT4J)

[Cure Duchenne](#) – Duchenne Muscular Dystrophy

[Cure GM1 Foundation](#) – GM1 Gangliosidosis

[Cure HHT](#) – Hereditary Hemorrhagic Telangiectasia

[Cure JM Foundation](#) – Juvenile Myositis, Juvenile Dermatomyositis, Amyopathic Dermatomyositis and Juvenile Polymyositis

[Cure RTD Foundation](#)– Riboflavin Transporter Deficiency Neuropathy

[Cure Sanfilippo Foundation](#) – Sanfilippo Syndrome

[Cure SMA](#) - Spinal Muscular Atrophy

[Cure SMA Foundation of India](#) - Spinal Muscular Atrophy

[Cure Surf1](#) – SURF1 Leigh Syndrome

[Cure TBM](#)– TracheoBronchoMalacia

[Cures Within Reach](#) – Multiple Rare Diseases

[Cure VCP Disease](#)

[CureGRIN](#) - GRIN

[CureLGMD2i Foundation](#)– Limb Girdle Muscular Dystrophy Type 2i

[CureSHANK](#) - Phelan-McDermid Syndrome and related SHANK3 disorders

[CureSPG50](#) - SPG50

[Curing Retinal Blindness Foundation](#) – Leber’s Congenital Amaurosis, Retinitis Pigmentosa

[Cushing’s Support and Research Foundation](#) – Cushing’s Syndrome

[Cutaneous Lymphoma Foundation](#) – Cutaneous Lymphoma Disease

[Cyclic Vomiting Syndrome Association](#) – Cyclic Vomiting Syndrome

[Cystic Fibrosis Research, Inc.](#) – Cystic Fibrosis

[Cystinosis Research Network](#) – Cystinosis

[Dana’s Angels Research Trust](#) – Niemann Pick Type C

[Danny’s Dose Alliance](#) – Rare Diseases, Chronic Illness

[Daphne’s Lamp](#)



[DDX3X Foundation](#) - General Support Organization  
[debra of America](#) - Epidermolysis Bullosa  
[Defeat MSA](#) - Multiple System Atrophy  
[Dent Disease Foundation](#) - Dent Disease  
[Determinence, Inc.](#) - Multiple Rare Diseases  
[DHPS Foundation](#)  
[DNA Checkup](#) - Recessive Genetic Disorders  
[Dravet Syndrome Foundation](#) - Dravet Syndrome  
[Dravet Syndrome Foundation Spain](#) - Dravet Syndrome  
[Dreamsickle Kids Foundation](#) - Sickle Cell Disease  
[Drépanovie- Association des Drépanocytaires de Côte D'Ivoire](#) - Sickle Cell Anemia  
[DRESS Syndrome Foundation](#) - Drug Reaction with Eosinophilia and Systemic Symptoms  
[Dup15q Alliance](#) - Chromosome 15q Duplication Syndrome  
[DYRK1A Community](#) - DYRK1A  
[Dystonia Medical Research Foundation](#) - Dystonia  
[EB Research Partnership](#) - Dystrophic Epidermolysis Bullosa  
[EBF3 HADDS Foundation](#) - EBF3-related HADDS  
[The EHE Foundation](#) - Epithelioid Hemangioendothelioma  
[EDMD International, Inc.](#) - Emery-Dreifuss Muscular Dystrophy  
[EDSers United](#) - Ehlers Danlos Syndrome  
[Ehlers-Danlos Network CARES Foundation](#) - Ehlers Danlos Syndrome  
[Ehlers-Danlos Society](#) - Global community focused on Ehlers-Danlos syndromes (EDS), hypermobility spectrum disorders (HSD) and related symptoms and conditions  
[Ehlers-Danlos Support Group of Greater Kansas City](#) - Ehlers Danlos Syndrome  
[Ehlers-Danlos Support Group of Jacksonville](#) - Ehlers Danlos Syndrome  
[Emerie Lee Foundation](#) - Childhood Blindness  
[Encephalitis Society](#) - Encephalitis  
[Endosalpingiosis Foundation Inc](#) - Endosalpingiosis  
[Emily's Entourage](#) - Cystic Fibrosis  
[Epidermoid Brain Tumor Society](#) - Epidermoid brain tumor  
[Epilepsy Alliance America](#) - Epilepsy, Seizure Disorders, Rare Epilepsies  
[Epilepsy Warriors Foundation](#) - Epilepsy  
[Erdheim-Chester Disease Global Alliance](#) - Erdheim-Chester Diseases  
[Ethan Lindberg Foundation](#) - Congenital Heart Disease  
[Ethan's Reason](#) - Batten Disease  
[EveryLife Foundation](#) - Multiple Rare Diseases  
[Fabry Australia](#) - Fabry Disease  
[Fabry Support and Information Group](#) - Fabry Disease  
[Facial Pain Association](#) - Trigeminal Neuralgia  
[FAM177A1](#) - Undiagnosed Community

[\*\*Familial Adenomatous Polyposis Foundation\*\*](#) – Familial Adenomatous Polyposis

[\*\*FamilieSCN2a Foundation\*\*](#) – SCN2a gene mutations

[\*\*FCS Foundation\*\*](#) – Familial Chylomicronemia Syndrome

[\*\*Feeding Tube Awareness Foundation\*\*](#) – Multiple Rare Diseases

[\*\*FH Foundation\*\*](#) – Familial Hypercholesterolemia

[\*\*Fibromuscular Dysplasia Society of America\*\*](#) – Fibromuscular Dysplasia

[\*\*Fibrous Dysplasia Foundation \(FDF\)\*\*](#) – McCune Albright syndrome, Fibrous dysplasia

[\*\*Fibrous Dysplasia Foundation FD/MAS Alliance\*\*](#) - Fibrous dysplasia/McCune Albright syndrome (FD/MAS Alliance)

[\*\*Fighting for Kaiden Foundation, Inc.\*\*](#) - Spinal Muscular Atrophy

[\*\*Fighting H.A.R.D. Foundation\*\*](#) – Rare Disease, Chronic Illness

[\*\*Findacure\*\*](#) – Multiple Rare Diseases

[\*\*FIRST: Foundation for Ichthyosis & Related Skin Types\*\*](#) – Skin Diseases, Ichthyosis

[\*\*FMD Chat\*\*](#) – Fibromuscular dysplasia

[\*\*FMF & AID Global Association\*\*](#) – Familial Mediterranean Fever & Autoinflammatory Diseases

[\*\*ForeBatten Foundation\*\*](#) - Juvenile Batten Disease

[\*\*Foundation for Angelman Syndrome Therapeutics\*\*](#) – Angelman Syndrome

[\*\*Foundation for Batten Hope\*\*](#) – Batten Hope

[\*\*Foundation for Mitochondrial Medicine\*\*](#)– Mitochondrial and Related Diseases

[\*\*Foundation for Prader-Willi Research\*\*](#) – Prader-Willi Syndrome, abnormal chromosome 15 (15q11-q13).

[\*\*Foundation for Sarcoidosis Research\*\*](#) – Sarcoidosis

[\*\*Foundation to Fight H-abc\*\*](#) - H-ABC

[\*\*FPIES Foundation\*\*](#) – Food Protein-Induced Enterocolitis Syndrome

[\*\*Friedreich's Ataxia Research Alliance\*\*](#) – Friedreich's Ataxia

[\*\*Fundacion Huntington Puerto Rico\*\*](#) - Huntington

[\*\*Fundación Red Sanfilippo\*\*](#) – Sanfilippo Syndrome / MPS III

[\*\*GACI Global\*\*](#) - Generalized Arterial Calcification of Infancy

[\*\*Garrett the Grand\*\*](#) – Batten Disease

[\*\*Gaucher Association of Australia & New Zealand\*\*](#) - Gaucher

[\*\*Gaucher Community Alliance\*\*](#) - Gaucher

[\*\*GBS/CIDP Foundation International\*\*](#) – Guillain-Barré Syndrome, Chronic Inflammatory Demyelinating Polyneuropathy

[\*\*Gene Giraffe Project\*\*](#) – Multiple Rare Diseases

[\*\*Genesic Nonprofit Organization, Incorporated\*\*](#) - Sickle Cell

[\*\*Genetic Alliance UK\*\*](#) – Multiple Rare Diseases

[\*\*Genetic Aortic Disorders Association\*\*](#) - Canada - Aortic Disorders

[\*\*Genetic Epilepsy Team Australia\*\*](#) - Genetic Epilepsies

[\*\*Glut1 Deficiency Foundation\*\*](#) - Glut1 Deficiency

[\*\*Global DARE Foundation\*\*](#) - Refsum Disease

[\*\*Global Hydranencephaly Foundation\*\*](#) – Hydranencephaly



[Global Liver Institute](#) - Liver Diseases

[Global Pediatrics MDS Initiative](#) - Global Pediatric MDS Initiative

[GNE Myopathy International](#) - GNE Myopathy, Hereditary Inclusion Body Myopathy , HIBM

[Grace Science Foundation](#) - NGLY1 Deficiency

[GRIN2B Foundation](#)

[Grupo De Estudos Doencas Raras](#) -Multiple Rare Diseases, Brazil

[Gwendolyn Strong Foundation](#) - Spinal Muscular Atrophy

[Hadley Hope Fund](#) - Niemann-Pick Type C disease

[HAE Canada](#) - Hereditary Angioedema

[Hairy Cell Leukemia Foundation](#) - Hairy Cell Leukemia

[Hannah's Hope Fund](#) - Giant Axonal Neuropathy

[Harmony 4 Hope](#) - Multiple Rare Diseases

[Harry's Fund](#) - Cri du Chat syndrome

[Haystack Project](#) - Multiple Rare Diseases

[HCU Network America](#) - Homocystinurias

[HD-Care](#) - Huntington's Disease

[Health Advocacy Summit](#)- Chronic Disease

[Help 4 HD International](#) - Huntington's disease

[Helping Hands for GAND, Inc.](#) - GATAD2B-associated neurodevelopmental disorder

[Hemophilia Foundation of Southern California](#)- Genetic Bleeding Disorders

[Hepatitis Foundation International](#) - Hepatitis

[Hereditary Angioedema Association](#) - Hereditary Angioedema

[Hereditary Hemorrhagic Telangiectasia Foundation International](#) - Hereditary Hemorrhagic Telangiectasia

[HESA - Hashimoto's Encephalopathy/SREAT & Seronegative Autoimmune Encephalitis Alliance](#) - Autoimmune Encephalitis, Hashimoto Encephalitis, Sero Negative Encephalitis

[Highway of Hope](#) - Multiple Rare Diseases

[Histiocytosis Association](#) - Histiocytic Disorders, Langerhans Cell Histiocytosis

[Hope4tylerandluke](#) - Med 23 gene defect

[Hope for Hypothalamic Hamartomas](#) - Hypothalamic Hamartomas

[Hope for Javier](#) - Duchenne Muscular Dystrophy

[HOPE for SCD](#) - Sickle Cell Disease

[Hope for Trisomy 13 & 18](#) - Trisomy 13 & 18 and related disorders, Edwards and Patau Syndrome.

[Hope Life International Gambia](#)- Pituitary, Adrenal, Achalasia, and Tuberculosis (TB)

[Hope Life Support & Empowerment \(HLSE\)- Nigeria](#) - Hope Life Support and Empowerment- Nigeria

[Hope Life Support and Empowerment \(HLSE\) Sierra Leone](#) - Hope Life Support and Empowerment - Sierra Leone

[HPS Network](#) - Hermansky-Pudlak Syndrome

[HSAN1E Society](#) - Hereditary Sensory and Autonomic Neuropathy, Type 1E

[HudsonAlpha Institute for Biotechnology](#)

[Humans with Knobloch Syndrome](#) - Knobloch Syndrome

[Huntington's Disease Society of America, Inc.](#) - Huntington's Disease

[Huntington's Disease Youth Organization \(HDYO\)](#)

[Hydrocephalus Association](#) - Hydrocephalus

[HyperIGM Foundation](#) - Hyper IGM Syndrome

[Hypoparathyroidism Association](#) - Hypoparathyroidism

[iCAN Research](#) - Multiple Rare Diseases

[iDefine](#) - Kleefstra Syndrome

[Idic15 Canada](#) - Chromosome 15q11.2-13.1 Duplication Syndrome

[IFAA International Foundation for Autoimmune and Autoinflammatory Arthritis](#)

[IHope Foundation](#) - Intracranial Hypertension

[Illinois Spina Bifida Association](#) - Spina Bifida

[INADcure](#) - Infantile Neuroaxonal Dystrophy

[Incontinentia Pigmenti International Foundation](#) - Incontinentia Pigmenti

[Indian Prader Willi Syndrome Association](#) - Prader Willi Syndrome

[IndoUSrare](#) - General Support Organization

[Instituto Vidas Raras \(Rare Lives Institute\)](#) - Brazilian umbrella organization for rare diseases

[Intermountain PKU and Allied Disorders Association](#) - Phenylketonuria and allied disorders

[International Cystinuria Foundation](#) - Cystinuria

[International Essential Tremor Foundation](#) - Essential Tremor

[International Firefighter Cancer Foundation, Inc.](#) - Multiple Rare Cancers

[International FOP Association](#) - Fibrodysplasia Ossificans Progressiva, Myositis Ossificans Progressiva

[International Foundation for CDKL5 Research](#)

[International FOXP1 Foundation](#) - FOXP1 gene mutation

[International Pemphigus & Pemphigoid Foundation](#) - Pemphigus, Pemphigoid

[International Prader-Willi Syndrome Organisation \(IPWSO\)](#) - Prader-Willi Syndrome

[International WAGR Syndrome Association](#) - WAGR Syndrome, 11p Deletion Syndrome

[IRF2BPL Foundation](#) - IRF2BPL Disorder

[ISMRD The International Advocate for Glycoprotein Storage Diseases](#) - Glycogen Storage Disease

[Jain Foundation](#) - Limb Girdle Muscular Dystrophy type 2B/R2/Mioyshi Myopathy/Dysferlinopathy

[Jamal's Helping Hands](#) - Multiple Chronic Illnesses

[Jamie's Dream Team](#) - Multiple Rare Diseases

[Jansen de Vries Syndrome Foundation](#) - Jansen de Vries Syndrome Foundation

[Jett Foundation](#) - Duchenne Muscular Dystrophy

[Jonah's Just Begun](#) - Sanfilippo Syndrome, Mucopolysaccharidoses (MPSIII)

[Jordan's Guardian Angels](#) - Jordan's Syndrome; Genetic misspelling of the genes PPP2R5D, PPP2R1A, PPP2R5C

[Joshua Frase Foundation](#) – Myotubular Myopathy

[Joshua Hellmann Foundation for Orphan Disease](#) – Multiple Rare Diseases, Hong Kong

[Joubert Syndrome & Related Disorders Foundation](#) – Joubert Syndrome

[Judson's Legacy](#) – Krabbe's Disease

[Kabuki Syndrome USA](#) –Kabuki syndrome

[KCNQ2 Cure](#) – KCNQ2

[KIF1A.org](#)

[KIF4A Foundation](#) - KIF4A Associated Neurodevelopmental Syndrome (KANS)

[Klippel-Feil Syndrome Freedom](#) – Klippel-Feil Syndrome

[Kneading Hope](#) – Multiple Rare Diseases

[KnowTheGlow](#) - Multiple Rare Diseases

[Koolen-de Vries Syndrome Foundation](#) - Koolen-de Vries Syndrome

[KrabbeConnect](#) – Krabbe

[Kruzn for a Kure Foundation](#) - Shimke Immuno Osseous Dysplasia

[K-T Support Group](#) – Klippel-Trenaunay Syndrome

[Kure for Kulas](#) – Spinal muscular atrophy (SMA)

[LAL Solace, Inc.](#) – Lysosomal Acid Lipase disease, LAL Deficiency, Wolman Disease

[Layla's Hope Foundation](#) – KCTD7 (Progressive Myoclonic Epilepsy Type 3)

[Lauren's Hope](#) – Hypomagnesemia with Secondary Hypocalcemia (HSH)

[Legg Calve Perthes Foundation](#) – Legg-Calve-Perthes Disease

[Leiomyosarcoma Direct Research Foundation](#) – Leiomyosarcoma

[LEMS Patients](#) – Lambert-Eaton Myasthenic Syndrome

[LGS Foundation](#) – Lennox-Gastaut Syndrome

[LHON Project](#) – Leber Hereditary Optic Neuropathy

[Life Branches NFP, Inc.](#) - Multiple Rare Diseases

[Li-Fraumeni Syndrome Association](#)– Li-Fraumeni Syndrome

[Lightning and Love Foundation](#) - Advocacy and research into THAP12 mutation related disease

[Lily's List](#) - Pediatric patients with a need for hospital-to-home health nursing

[Little Zebra Fund](#) - Undiagnosed & General Support Organization

[Lipodystrophy United](#) – Lipodystrophy

[Lipoprotein\(a\) Foundation](#) – High Lipoprotein(a)

[Little Hercules Foundation](#)– Duchenne Muscular Dystrophy

[Little Miss Hannah Foundation](#) – Multiple Pediatric Rare Diseases

[Littlest Tumor Foundation](#) - Neurofibromatosis

[Living in the Light of Rare Diseases](#) - Multiple Rare Diseases

[Live4TheCure](#) – All Rare Diseases

[Living LFS](#) – Li-Fraumeni Syndrome

[LouLou Foundation](#) – CDKL5 Deficiency Disorder

[Love Never Sinks](#) - General Support Organization  
[Low Syndrome Association](#) - Low Syndrome  
[LSD Society](#) - LSD, Pakistan  
[Luka the Lion Foundation](#) - Multiple Pediatric Rare Diseases  
[Lymphangiomatosis & Gorham's Disease Alliance](#) - Lymphangiomatosis, Gorham's Disease  
[Lynch Syndrome International](#) - Lynch Syndrome  
[Lysosomal Diseases New Zealand](#)  
[Lysosomal Storage Disorders Support Society](#) - Lysosomal Diseases, India  
[M-CM Network](#) - Macrocephaly-capillary malformation  
[Mackenzie's Mission](#) - Amyloidosis  
[MacPac Foundation](#) - Hypomyelination with brainstem and spinal cord involvement and leg spasticity (HBSL)  
[Malan Syndrome Foundation](#) - Malan Syndrome  
[Marrow For Life, Inc.](#) - Stem Cell Treatments for Rare Diseases  
[Mason Shaffer Foundation](#) - Malignant Infantile Osteopetrosis  
[Massachusetts Biotechnology Council](#) - Multiple Rare Diseases  
[Mast Cell Hope, Inc.](#) - Mast Cell Disease  
[MCT8-AHDS Foundation](#) - MCT8 Deficiency aka Allen Herndon Dudley Syndrome  
[MECP2 Duplication Foundation](#) - MECP2 Duplication Syndrome  
[Meg's Miracles](#) - Childhood / Pediatric and Adult Onset SPS  
[MEPAN.org](#) - MEPAN Syndrome  
[Metaplastic Breast Cancer Global Alliance](#)  
[Meningitis Foundation of America](#)  
[MHE Coalition](#) - Multiple Hereditary Exostoses, Multiple Osteochondromatosis, Hereditary Multiple Exostoses  
[Mickie's Miracles](#) - Infantile Spasms, Pediatric Epilepsy  
[Miles for CF](#) - Cystic Fibrosis  
[Milo's Journey Foundation](#) - Undiagnosed  
[Minutes Matter](#) - Medium-chain acyl-CoA dehydrogenase deficiency (MCADD)  
[Miracle Flights](#) - All Rare Diseases  
[Miracle For Madison & Friends](#) - Spinal Muscular Atrophy  
[Mississippi Metabolics Foundation](#) - Rare Genetic Metabolic Disorders / IEMs  
[Mission: Cure](#)  
[Mission Massimo Foundation](#) - Leukodystrophy  
[MitoAction](#) - Mitochondrial Disease  
[MLD Foundation](#) - Metachromatic Leukodystrophy  
[Moebius Syndrome Foundation](#) - China Doll Syndrome, Moebius Syndrome  
[Mommies of Miracles](#) - Multiple Pediatric Rare Diseases  
[Mo Songs for Kerry](#) - Li Fraumeni Syndrome  
[Moyamoya Foundation Co](#) - Moyamoya Disease

[MSA-AMS and World MSA Day](#)

[MSA NJ](#)– Multiple System Atrophy

[MSUD Family Support Group](#)

[MTM-CNM Family Connection, Inc.](#) – Myotubular and Centronuclear Myopathy

[Muscular Dystrophy Family Foundation](#) Muscular Dystrophy, Neuromuscular Diseases

[Muscular Dystrophy Society Kenya](#) – Muscular Dystrophy

[My PKU Awareness Foundation](#) - PKU

[Myasthenia Gravis Hope Foundation](#)– Myasthenia Gravis

[Myhre Syndrome Foundation](#) - Myhre Syndrome (SMAD4 Mutation)

[Myositis Support and Understanding Association](#) - Idiopathic Inflammatory Myopathies, Myositis

[Myotonic Dystrophy Foundation](#) – Myotonic Dystrophy

[Myrovlytis Trust](#) – Birt Hogg Dube Syndrome

[Narcolepsy Network](#) – Narcolepsy

[National Ataxia Foundation](#) - Ataxia

[National Foundation for Ectodermal Dysplasias](#) – Ectodermal Dysplasias

[National Fragile X Foundation](#) - National Fragile X Foundation

[National Gaucher Foundation](#) – Gaucher Disease

[National Hemophilia Foundation](#) - Hemophilia

[National LeioMyoSarcoma Foundation](#) – LeioMyoSarcoma

[National Lymphedema Network](#) – Lymphedema-distichiasis syndrome, Hereditary lymphedema type II & other lymphedemas

[National Marfan Foundation](#) – Marfan Syndrome

[National MPS Society](#) – Mucopolysaccharidoses

[National Niemann-Pick Disease Foundation](#) – Niemann Pick Disease

[National Organization of Disorders of the Corpus Callosum \(NODCC\)](#) - Disorders of the Corpus Callosum

[National PKU Alliance](#) – Phenylketonuria

[National PKU News](#) – Phenylketonuria

[National Spasmodic Dysphonia Association](#) – Spasmodic Dysphonia, Laryngeal Dystonia

[National Stem Cell Foundation](#)

[National Tay-Sachs & Allied Disease Association](#) – Tay-Sachs, Canavan, Sandhoff, GM1 & related diseases

[NBIA Disorders Association](#) – Aceruloplasminemia, Neurodegeneration w/ Brain Iron Accumulation disorders, Hallervorden-Spatz Syndrome

[NEHI Research Foundation](#)– Neuroendocrine cell hyperplasia of infancy

[NephCure Kidney International](#) - Rare Forms of Nephrotic Syndrome

[Neurofibromatosis Midwest](#) - Neurofibromatosis and Schwannomatosis

[Neurofibromatosis Network](#) – Neurofibromatosis Disease

[Neuromuscular Disease Foundation](#) – Neuromuscular Disease

[Nevus Outreach, Inc.](#) – Congenital Melanocytic Nevi and Related Disorders

[Newborn Coalition Foundation](#) – Newborn Screening

[New Zealand Organisation for Rare Disorders](#) – Multiple Rare Diseases

[New Zealand Pompe Network](#)

[Next Step Fund](#) – Life Threatening Diseases

[NF2BioSolutions](#) – Neurofibromatosis Type 2

[NGLY1 Foundation](#)– NGLY1 deficiency, N-glycanase deficiency

[Nicholas Volker One In A Billion Foundation](#)

[Niemann Pick Research Foundation](#) – Neimann Pick Disease

[Noah’s Hope-Hope 4 Bridget Foundation](#) – Batten Disease

[Nonsense Mutations Foundation](#) - Nonsense Gene Mutations

[Nontuberculous Mycobacteria Info & Research](#) – Pulmonary Nontuberculous Mycobacterial

[Noonan Syndrome Foundation](#) – Noonan Syndrome

Noonan Syndrome Support Network – Noonan Syndrome

[NR2F1 Foundation](#) - Bosch Boonstra Schaaf optic atrophy syndrome

[NUBPL Foundation](#)

[One Rare](#) - Serving young adults with rare and chronic conditions

[Organic Acidemia Association](#) – Organic Acidemia

[Organization for Rare Diseases India](#) – Multiple Rare Diseases

[Orphan Disease Center](#)– Multiple Rare Diseases

[Osteopetrosis Society](#) – Osteopetrosis

[Ostomy 2-1-1 Inc.](#) - Ostomy, Short Gut, Short Bowel

[Our Odyssey](#) - All rare and chronic conditions for young adults 18-35

[Oxalosis and Hyperoxaluria Foundation](#) – Hyperoxaluria, Oxalosis

[Pachyonychia Congenita Project](#) – Pachyonychia Congenita

[PANDAS Network.org](#) – Pediatric Autoimmune Neuropsychiatric Disorders Associated with Streptococcal Infections

Parent/Consumer Advocate for MS

[Parent Project Muscular Dystrophy](#) – Duchenne Muscular Dystrophy, Becker Muscular Dystrophy

[Pathways for Rare and Orphan Studies](#) (PROS) – Multiple Rare Diseases

[Pathways To Trust](#) - General Support Organization, current focus Sickle Cell Disease

[Patient Advocate Foundation](#) - General Support Organization

[PBCers Organization](#)

[PCDH19 Alliance](#) – PCDH19 Epilepsy

[Pediatric Brain Foundation](#) – Pediatric Neurologic Disorders

[Pediatric Hydrocephalus Foundation](#) – Hydrocephalus

[Perthes Kids Foundation](#) – Legg-Calvé-Perthes Disease

[Pericarditis Alliance](#) - Pericarditis Alliance

[Petronille Healthy Society](#) - Sickle Cell Anemia



[Phelan-McDermid Syndrome Foundation](#) – 22Q13 Deletion Syndrome, Phelan-McDermid Syndrome

[Pheo Para Alliance](#) – Pheochromocytoma & Paraganglioma Diseases

[Pheo Para Project](#) – Pheochromocytoma & Paraganglioma

[PHIP Kids \(Chung-Jansen Syndrome\)](#) - Chung-Jansen Syndrome

[Pitt Hopkins Research Foundation](#) – Pitt Hopkins

[Pituitary Network Association](#) – Pituitary Disorders

[PKD Foundation](#) – Polycystic Kidney Disease

[PKS Kids](#) – Pallister-Killian Syndrome

[PKS Kids Italia Onlus](#) – Pallister-Killian Syndrome

[Platelet Disorder Support Association](#) – Immune Thrombocytopenia

[PMG Awareness Organization, Inc.](#) – Polymicrogyria

[PMP Pals Network](#) - Pseudomyxoma Peritonei

[Pompe Alliance](#) - Pompe Disease

[Pompe Foundation, India](#) – Pompe Disease

[Pompe Warrior Foundation](#) - Pompe Disease

[Potocki-Lupski Syndrome Outreach Foundation, Inc](#) – Potocki-Lupski Syndrome

[Prader-Willi California Foundation](#) – Prader-Willi Syndrome

[Prader-Willi Syndrome Association](#) | USA – Prader-Willi Syndrome

[PRISMS, Inc.](#) – Smith-Magenis Syndrome

[Project 8P](#) – Chromosome 8p

[Project Alive](#) - Hunter Syndrome or Mucopolysaccharidosis/MPS II

[PROS Foundation](#)

[Progeria Research Foundation](#) – Hutchinson-Gifford Progeria Syndrome

[Progressive Familial Intrahepatic Cholestasis Advocacy and Resource Network, Inc.](#)  
– Progressive Familial Intrahepatic Cholestasis

[PRP Alliance](#) – Pityriasis Rubra Pilaris

[PSC Community](#) – Primary Sclerosing Cholangitis

[PSC Partners](#) - Primary Sclerosing Cholangitis

[PTEN Hamartoma Tumor Syndrome Foundation](#)

[PTen World](#) – Cowden’s Syndrome

[Pulmonary Hypertension Association](#) – Pulmonary Hypertension

[PURA Syndrome Foundation](#) – PURA Syndrome

[Putting Rare Diseases Patients First!](#) - All Rare Diseases

[PVNH Support and Awareness](#) – Periventricular Nodular Heterotopia

[Pyruvate Kinase Deficiency](#) - Pyruvate Kinase Deficiency

[RACC- UK](#) – Periodic Fever Syndromes

[Raiden Science Foundation](#) - UBA5 Gene Mutation

[Raising Hope International Friends](#) - Sickle Cell Disease

[RareKC](#) – All Rare Diseases

[Rare & Undiagnosed Network](#) – Multiple Rare Diseases, Undiagnosed

[RareABILITY](#) - Hosts empowerment workshops and crafting activities for people of varying abilities

[Rare Advocacy Movement](#) - All Rare Diseases

[Rare Disease Awareness Men's Group](#) - Multiple Rare Diseases

[Rare Disease Foundation](#) – Multiple Rare Diseases, Canada

[Rare Disease Ghana Initiative](#)– Multiple Rare Diseases, Ghana

[Rare disease Innovations Institute](#) - Multiple Rare Diseases

[Rare Disease Kenya](#) – Multiple Rare Diseases, Kenya

[Rare Disease UK](#) – Multiple Rare Diseases

[Rare Disease Research and Support \(RDRS\) at Calvin University](#) - All Rare Diseases

[Rare Genomics Institute](#) – Multiple Rare Diseases

[Rare Kids Network, Inc.](#) - Pediatric Rare Diseases

[Rare New England](#) – Multiple Rare Diseases

[Rare Science](#) – Multiple Rare Diseases

[Rare Support](#) – Rare United Network

[Rare Trait Hope Fund](#) - Aspartylglucosaminuria

[Rare Village Foundation](#) - General Support Organization for several rare disorders

[RARE-X](#) - Rare Disease Data Collection Platform (Diagnosed and Undiagnosed)

[Raríssimas – Associação Nacional de Deficiências Mentais e Raras](#) – Multiple Rare Diseases, Portugal

[RASopathies Network USA](#) – RASopathy Syndromes

[Raymond A. Wood Foundation](#) - Hypothalamic-Pituitary Brain Tumor Survivors

[RecurrentMeningitis Association](#) – Recurrent Meningitis

[Recurrent Respiratory Papillomatosis Foundation](#) - Recurrent Respiratory Papillomatosis, Pulmonary Recurrent Respiratory Papillomatosis, Laryngeal Papilloma, Glottal Papilloma, HPV

[Redmoon Project, Inc](#) - Lifting Sickle Cell Warriors Across The World

[Reflex Sympathetic Dystrophy Syndrome Association \(RSDSA\)](#) – Complex Regional Pain Syndrome

[Relapsing Polychondritis Awareness and Support Foundation](#) – Relapsing Polychondritis

[Remember the Girls](#) – X-linked Recessive Diseases

[Rettsyndrome.org](#) – Rett Syndrome

[Rhode Island Ehlers Danlos Syndrome Awareness and Support Group](#)– Ehlers Danlos Syndrome

[Riaan Research Initiative](#) - Cockayne Syndrome

[Ring 14 USA](#) – Chromosome 14 disorders

[RUNX1 Research Program](#) - RUNX1-Familial Platelet Disorder

[Ryan's Challenge](#) - Vascular Ehlers Danlos Syndrome

[Ryan Wersten MIOP Foundation](#) – Malignant Infantile Osteopetrosis

[Salla Treatment and Research Foundation](#) – Salla Disease

[Sanfilippo Children's Foundation](#) – SanFilippo

[Sanford Research](#) – Multiple Rare Diseases

[Sarcoidosis of Long Island](#) – Neurosarcoidosis, Sarcoidosis

[Sarcoma Foundation of America](#) – Sarcoma

[SATB2 Gene Foundation](#)– SATB2 Associated Syndrome

[Save Sight Now](#) - Usher Syndrome Type 1B

[SBS Cure Project](#) – Short Bowel Syndrome/Intestinal Failure

[SCAD Alliance](#) – Spontaneous Coronary Artery Dissection

[Scheuermann's Disease Fund](#) - Scheuermann's Disease Fund

[SCN2A Australia](#) - SCN2A

[SETBP1 Society](#) – SETBP1

[SHANK 2 Foundation](#) - SHANK 2 Disorders

[Shwachman-Diamond Syndrome Alliance](#) - Shwachman-Diamond Syndrome

[Shwachman-Diamond Syndrome Foundation](#) – Shwachman-Diamond Syndrome

[Sick Cells](#) - Sickle Cell Disease

[Sickle Cell Red](#) - Sickle Cell Disease

[Sickle Cell Association of Texas Mark Thomas Foundation](#) - Sickle Cell General Support Organization

[Sickle Cell 101](#) – Sickle Cell Disease

[Sickle Cell and Young Stroke Survivors](#) - Sickle Cell Disease and Others

[Sickle Cell Consortium](#) – Sickle Cell Disease

[Sing Me a Story](#) - General Support

[Sisters' Hope Foundation](#) - Adult Onset Leukoencephalopathy with Axonal Spheroids and Pigmented Glia

[SLC6A1 Connect](#) – SLC6A1 (GAT1)

[SMC1A Foundation](#) - SMC1A Loss-of-function Epilepsy

[Smith-Kingsmore Syndrome](#) – Smith-Kingsmore Syndrome

[SMA - Bhutan](#) - Spinal Muscular Atrophy - Bhutan

[SMS Research Foundation](#) – Smith-Magenis Syndrome

[Sofia Sees Hope, Inc.](#)

[Soft Bones Inc.](#) – Hypophosphatasia

[Soft Bones Canada](#) – Hypophosphatasia

[Sophies Hope Foundation Inc](#) - Glycogen Storage Disease Type 1b

[Spastic Paraplegia Foundation](#) - Primary Lateral Sclerosis (PLS) and Hereditary Spastic paraplegia

[Spinal CSF Leak Foundation](#) – Cerebrospinal Fluid

[SPG15 Research Foundation](#) - SPG15

[SSADH Association](#)– Succinic Semialdehyde Dehydrogenase Deficiency

[St. Joseph's Children's Hospital](#), New Jersey - Multiple Rare Diseases

[Step by Stone Association](#)– Multiple Rare Diseases, Kenya

[Stone Soup Group](#) - All Rare Diseases

[Stop ALD Foundation](#) – X-linked adrenoleukodystrophy  
[Stories Matter Foundation](#) - General Support Organization  
[Stronger Than Sarcoidosis](#) - Sarcoidosis  
[STXBP1 Disorders/Foundation](#)  
[Super Kids: Mission Zero Inc.](#) – Epilepsy  
[Super T's Mast Cell Foundation](#) - Mast Cell Activation Disorder  
[Supporters of Families with Sickle Cell Disease, Inc.](#) – Sickle Cell Disease  
[Support Organization for Trisomy 18, 13 & Related Diseases](#) – Trisomy 13 Disorder, Trisomy 18

[Sweet Nectar Society](#) – Multiple Rare Diseases  
[Syndrome Without a Name \(SWAN\) Australia](#) - Undiagnosed  
[Syndromes Without A Name USA](#) – Undiagnosed  
[SynGap Research Fund](#) - SynGap  
[Tango2 Research Foundation](#) - Tango2  
[Taylor's Tale](#) – Batten Disease  
[T.E.A.M. 4 Travis](#) - Isolated Congenital Asplenia  
[Team Impact](#)  
[Team Jeffery for PF](#)  
[Team Joseph](#) - Duchenne  
[Team Telomere](#) - Dyskeratosis Congenita  
[Team Titin](#) – Titinopathies  
[Teddington Trust](#) – Xeroderma Pigmentosum  
[Tennessee PKU Foundation](#) - Phenylketonuria (PKU)  
[The 22q Family Foundation](#) – 22q11.2 deletion, DiGeorge Syndrome, VCFS  
[The AADC Research Trust](#)– Aromatic Amino Acid Decarboxylase Deficiency (AADC)  
[The Aarskog Foundation](#)– Aarskog-Scott Syndrome  
[The Akari Foundation](#) - Duchenne Muscular Dystrophy  
[The APS Type 1 Foundation](#)– Autoimmune Polyglandular Syndrome Type 1  
[The ARM Foundation](#) - The ARM Foundation  
[The Avery's Fight Foundation](#) - Homozygous Familial Hypercholesterolemia (HoFH)  
[The Avalon Foundation](#) - Hypophosphatasia, growing into General Support Organization  
[The Bonnell Foundation](#) - Cystic Fibrosis  
[The Boston International Turner Syndrome Summit](#) - Turner Syndrome  
[The Bow Foundation](#)– GNAO1  
[The Brain Recovery Project](#) – Multiple Rare Diseases  
[The Burning Limb Foundation](#)  
[The Calliope Joy Foundation](#)– Leukodystrophy  
[The Champ Foundation](#) - Single large scale mitochondrial DNA deletion syndromes (Pearson syndrome)  
[The CHARGE Syndrome Foundation](#)– CHARGE Syndrome

[\*\*The Charles E. Holman Morgellons Disease Foundation\*\*](#) – Morgellons Disease

[\*\*The Chromosome 18 Registry & Research Society\*\*](#) – Chromosome 18 disorders

[\*\*The Cilly Cell Project\*\*](#) – Sickle Cell Disease

[\*\*The Coalition Against Pediatric Pain\*\*](#) – Multiple Pediatric Rare Diseases

[\*\*The Conley Cushing's Disease Fund\*\*](#) – Cushing's Disease

[\*\*The COPD Foundation\*\*](#) – Chronic Obstructive Pulmonary Disease

[\*\*The Cure Blau Syndrome Foundation\*\*](#) - Blau Syndrome

[\*\*The Cure Tay-Sachs Foundation\*\*](#)

[\*\*The Cute Syndrome\*\*](#) – PCDH19 Epilepsy and SCN8A Epilepsy

[\*\*The Dent Disease Foundation\*\*](#) - Dent Disease

[\*\*The DRESS Syndrome Foundation\*\*](#) - Drug Reaction with Eosinophilia and Systemic Symptoms

[\*\*The FAVA Foundation\*\*](#) – Fibro Adipose Vascular Anomaly

[\*\*The Genesis Foundation for Children\*\*](#) - Multiple Rare Diseases

[\*\*The Global Foundation for Peroxisomal Disorders\*\*](#) –Peroxisome Biogenesis Disorder, Zellweger Spectrum Disorder

[\*\*The Hypersomnia Foundation\*\*](#)– Idiopathic Hypersomnia

[\*\*The Jansen's Foundation\*\*](#)– Jansen Type Metaphyseal Chondrodysplasia

[\*\*The LAMFoundation\*\*](#) – Lymphangiomyomatosis (LAM)

[\*\*The Life Raft Group\*\*](#) – Gastrointestinal Stromal Tumor (GIST)

[\*\*The Marfan Foundation\*\*](#) – Marfan Syndrome

[\*\*The Mastocytosis Society Inc\*\*](#) – Mastocytosis and Mast Cell Activation Disease

[\*\*The Mastocytosis Society Inc Southern California Support Group\*\*](#)

[\*\*The Menkes Foundation\*\*](#) - General Support Organization

[\*\*The Metabolic Foundation\*\*](#) – Metabolic Disorders

[\*\*The Mitchell Thorp Foundation\*\*](#) - General Support Organization

[\*\*The MOG Project\*\*](#) - Myelin Oligodendrocyte Glycoprotein Antibody Disease

[\*\*The Morgan Leary Vaughan Fund\*\*](#)

[\*\*The Mowat-Wilson Syndrome Foundation\*\*](#) – Mowat-Wilson Syndrome Foundation

[\*\*The Myositis Association\*\*](#) – Myositis

[\*\*The National Gaucher Foundation of Canada\*\*](#) - Gaucher

[\*\*The NICER Foundation\*\*](#) – Narcolepsy

[\*\*The Network of Tyrosinemia Advocates\*\*](#) – Tyrosinemia

[\*\*The Rare Genomes Project at the Broad Institute of MIT and Harvard\*\*](#)– Multiple Rare Diseases

[\*\*The Sickle Cell Foundation of Tennessee\*\*](#) - Sickle Cell

[\*\*The Snyder-Robinson Foundation\*\*](#) – Snyder-Robinson Syndrome

[\*\*The Speak Foundation\*\*](#) – Neuromuscular Diseases

[\*\*The Sturge-Weber Foundation\*\*](#) – Sturge-Weber Syndrome

[\*\*The Sumaira Foundation for NMO\*\*](#) - Neuromyelitis Optica Spectrum Disorder

[\*\*The TESS Foundation\*\*](#) – SLC13A5

[The Ultra Rare Disease Disorders and Disabilities Foundation](#) – Multiple Rare Diseases

[The Vision of Children Foundation](#) – Genetic Vision Disorders

[The WeHeal Foundation](#) - All Rare Diseases

[The Wiedemann-Steiner Syndrome Foundation](#) – Weidemann-Steiner Syndrome

[Theo's Village: The TBCK Foundation](#) - TBCK

[ThinkGenetic Foundation](#)– Multiple Rare Diseases

[Thisbe and Noah Scott Foundation, Inc.](#) – Pediatric Neurologic Disorders

[Timothy Syndrome Alliance](#) – Timothy Syndrome and CACNA1c mutations

[Translational Genomics Research Institute](#) – Multiple Rare Diseases

[Translational Pulmonary and Immunology Research Center](#) – Multiple Rare Diseases

[Transplant Unwrapped](#) - Short Bowel Syndrome, Chronic Intestinal Pseudo Obstruction, Intestinal Failure

[Traveling Awareness Bears](#) – Multiple Rare Diseases

[Trisomy 18 Foundation](#) – Trisomy 18, Edwards Syndrome

[TUBB3 Foundation](#)– TUBB3 Gene Mutations

[Tuberous Sclerosis Alliance](#)– Tuberous Sclerosis

[Tuberous Sclerosis Australia](#)

[Turner Syndrome Global Alliance](#) – Turner Syndrome

[Turner Syndrome Society of the United States](#) – Turner Syndrome

[U.R. Our Hope](#) – Undiagnosed

[United Leukodystrophy Foundation](#) – Leukodystrophies

[United Mitochondrial Disease Foundation](#) – Mitochondrial Encephalomyopathy

[Unique - Rare Chromosome Disorder Support Group](#) - Rare Chromosome and Single Gene Disorders

[University of Washington/Cystic Fibrosis Reproductive and Sexual Health Collaborative](#)

[Uplifting Athletes](#) - Umbrella Organization

[Usher 1F Collaborative, Inc](#) – Usher Syndrome Type 1F

[Usher Syndrome Coalition](#) – Usher Syndrome

[USP7 Families](#) – USP7 Related Diseases

[USTMA Consortium](#) - Thrombotic Microangiopathies

[Utah Rare](#)– Multiple Rare Diseases

[Vancouver Acromegaly Support Group](#) – Gigantism, Acromegaly

[VHL Alliance](#) - VHL

[Vincent Gaynor & Sophia's Cure Foundation](#) – Spinal Muscular Atrophy

[Vision for Tomorrow Foundation](#) - Aniridia, Albinism

[VWM Families Foundation, Inc](#) – Vanishing White Matter Disease VWM/CACH

[Wake Up Narcolepsy](#) – Narcolepsy

[WeCareJourney](#) - Spinal Muscular Atrophy (SMA)

[White Sutton Syndrome Foundation](#) – White Sutton Syndrome (WHSUS)

[Wilhelm Foundation](#) – Undiagnosed



[William E Proudford Sickle Cell Fund](#) – Sickle Cell Disease  
[Williams Syndrome Association](#) – Williams Syndrome  
[Wilson Disease Association](#) - Wilsons Disease  
[Wiskott Aldrich Foundation](#) – Wiskott-Aldrich Syndrome  
[Wobbly Feet Foundation](#) – Ataxia-Telangiectasia  
[WonderFIL Smiles](#) - General Support Organization  
[Wylder Nation](#) – Lysosomal Diseases  
[Xia-Gibbs Society, Inc.](#) - Xia-Gibbs Syndrome  
[XLH Network](#) – X-Linked Hypophosphatemia  
[XP Family Support Group](#) – Xeroderma Pigmentosum  
[Zoe's Story](#) – CRMO/CNO  
[ZTTK SON-Shine Foundation](#) - Zhu-Tokita-Takenouchi-Kim Syndrome