



Connecting

Minds

Changing

Lives



Message From Leadership



MARK RUDOLPH
President



JANET L. HIESHETTER
Executive Director

In 2023, the Dystonia Medical Research Foundation made connections. We brought together the best and brightest minds to advance dystonia research and gathered the most dedicated support leaders to share ideas to help individuals and families with dystonia. We also connected the dystonia community with one another through online and in person events to fight the feeling of isolation that can occur with this life-changing movement disorder.

Your support and trust in the DMRF made these connections possible, driving hope and changing lives. Examples of these critical connections can be seen throughout this Annual Report, but we'll spotlight a few here.

The much-anticipated Samuel Belzberg 6th International Dystonia Symposium (IDS6), held in June in Dublin, Ireland, drew world-renown scientists from a broad spectrum of dystonia research. Researchers from 32 countries gathered for three days to discuss current research and debate new areas to pursue going forward in all aspects of the dystonia field.

DMRF co-founder Frances Belzberg kicked off the Symposium by thanking the researchers for their work and the progress that has been made in our understanding of dystonia. She also reminded them that people like her daughter are still waiting for scientific breakthroughs and answers. DMRF's quest to improve dystonia treatments and ultimately find a cure is not finished, and we need continued collaboration between the scientific and patient communities.

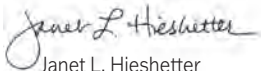
In March of 2023, the DMRF connected top minds in another area of dystonia by hosting a virtual meeting on the non-motor symptoms of dystonia, bringing much needed attention to this important aspect of the movement disorder. It is extraordinarily important that we examine connections between dystonia and things like anxiety, social anxiety, and depression. This meeting and the work of DMRF's Mental Health Programming Committee reflect the Foundation's commitment to shining a light on these very real concerns of our community.

DMRF's support network also promotes critical connections in the dystonia community. We hosted more than 135 support group meetings in 2023, led by volunteer support leaders who are living with dystonia and understand its challenges and the isolation that a diagnosis can bring. To encourage and educate support leaders, DMRF hosted two virtual support leader gatherings where leaders tapped into the knowledge and experiences of other support leaders from around the country.

All that we achieved in 2023 was made possible because of you. Thank you for supporting the DMRF's mission with your financial support, volunteer efforts and participation in DMRF programs. Together, we will continue connecting minds to change the lives of those who turn to the DMRF for hope through research, accurate information, support and community.



Mark Rudolph
President



Janet L. Hieshetter
Executive Director

Mission

The mission of the Dystonia Medical Research Foundation (DMRF) is to advance research for more effective treatments and ultimately a cure, to promote awareness and education, and to support the well-being of affected individuals and families.

About Dystonia

Dystonia is a neurological disorder that causes excessive, involuntary muscle contractions. These muscle contractions force the body and limbs into abnormal movements and postures, making it difficult for individuals to control their movements. Dystonia can affect virtually any region of the body including the face, jaw, eyelids, neck, vocal cords, torso, limbs, hands, and feet. Dystonia may affect a single body area or be generalized throughout multiple muscle groups. Dystonia affects men, women, and children of all ages and backgrounds. Estimates suggest no fewer than 250,000 people in the United States are affected.

Dystonia causes varying degrees of disability and pain, from mild to severe. Although treatments exist, dystonia remains a chronic disorder for which there is not yet a cure.

Samuel Belzberg 6th International Dystonia Symposium Showcases Latest Research

The Samuel Belzberg 6th International Dystonia Symposium drew top dystonia researchers from 32 countries, including 68 faculty members and close to 350 attendees. The Symposium also showcased 105 scientific posters presenting the latest in dystonia research.

After 12 years of waiting, five years of preparations and a rescheduling due to the global pandemic, the DMRF finally was able to gather dystonia researchers from 32 countries in Dublin, Ireland for the Samuel Belzberg 6th International Dystonia Symposium (IDS6) in June of 2023.

A showcase of progress and advancement in all aspects of the dystonia field, the symposium featured three days of meticulously planned sessions, carefully negotiated topics, and participation of essentially everyone deeply involved in dystonia research.

With IDS6, our goal was to extend knowledge beyond current boundaries, highlight important scientific advances in the field, stimulate discussion within and across disciplines, and push forward into a deeper understanding of the disorder and how to optimize treatment for those who are affected.

Nearly 350 attendees participated in IDS6, which showcased 105 scientific posters presenting the latest dystonia research. Talks ranged from fundamental summaries of clinical aspects of the dystonias to the very latest developments in the scientific understanding of dystonia thanks to phenomenal progress in genetics, neurophysiology, animal modelling, and other areas of dystonia neuroscience. IDS6 was a huge success and represents the latest chapter in a definitive series of international dystonia meetings organized by the DMRF since 1975.

The IDS6 Scientific Program Committee was co-chaired by H. A. (Buz) Jinnah, MD, PhD, and Antonio Pisani, MD, PhD, and included members from all over the world. The DMRF also supported 30 travel scholarships for promising young investigators to attend this outstanding international symposium.

“As an aspiring principal investigator in the field of dystonia research, the Samuel Belzberg 6th International Dystonia Symposium provided me with a rare opportunity to learn about where we are and where we are going in dystonia research, to network with leading researchers in the field, to present my work to an international audience and to inspire me with new ideas to consider in my own research.”

*Alejandro Rey Hipolito, PhD Candidate,
Baylor College of Medicine
(DMRF travel scholarship recipient)*



“Dystonia advocates are fully engaged and work tirelessly to support our legislative agenda to benefit the dystonia community. Because of their many efforts, dystonia continues to be included on the list of eligible conditions through the Department of Defense Peer Review Medical Research Program. This has been an important resource for dystonia researchers with more than \$28 million awarded since 2010.”

*Carole Rawson,
DMRF Vice President
of Public Policy*



Dystonia Advocacy Drives Awareness

The Dystonia Advocacy Network (DAN) advocates for all persons affected by dystonia and supports a legislative and policy agenda that meets the needs of the dystonia community.



In 2023, the DMRF continued to provide staff support to the collaborative efforts of DAN, which includes the DMRF, Benign Essential Blepharospasm Research Foundation, Dysphonia International, and the National Spasmodic Torticollis Association. Dystonia advocates met virtually with Congressional offices about the needs of the dystonia community and worked to educate elected officials about dystonia and how it changes the lives of those affected. Advocates were once again successful in having dystonia included on the list of eligible conditions through the Department of Defense Peer Review Medical Research Program. This program has been an important funding source for dystonia researchers with more than \$28 million awarded since dystonia was first included in 2010.

Dystonia Coalition

The DMRF continues to serve as the administrative center at no cost for the **Dystonia Coalition**, a collaboration of medical researchers and patient advocacy groups supported by the Office of Rare Diseases

Research in the National Center for Advancing Translational Sciences and the National Institute of Neurological Disorders & Stroke (NINDS) at the National Institutes of Health. The mission is to advance the pace of clinical and translational research in the dystonias to find better treatments and a cure.

The Coalition works to develop a fuller understanding of the many features of dystonia and how they change over time, to develop validated diagnostic strategies and rating tools for diagnosis and monitoring patients in clinical trials, and to establish a biorepository where blood and other samples can be stored and distributed for research. The work of the Coalition continues to stimulate clinical trials, promote dystonia education and awareness, and produce new articles and research.

Dystonia Brain Collective

The **Dystonia Brain Collective** promotes brain donation registration to further research and provide investigators with access to critically important brain samples.

The DMRF continues to serve as the administrative center and reviews and processes tissue requests from qualified researchers for approved dystonia related research. The Dystonia Brain collective has 884 registered brain donors in its database currently. Forty-seven brains have been recovered and are stored at the Harvard Brain Tissue Resource Center; one brain was recovered in 2023.

The Global Dystonia Registry

The DMRF continues to provide the operations and administration of **The Global Dystonia Registry** (GDR). GDR supports future dystonia studies through the confidential collection of patient data and distribution to dystonia investigators. To date, more than

6,600 people have registered worldwide.

Dystonia Journal Publishes Premier Research

The DMRF's journal, Dystonia, published 28 articles in 2023 with a total of 40 articles since its inception two years ago. Dystonia aims to be the leading journal in the field by publishing premier research on all basic, clinical, and translational aspects of dystonia, providing a centralized publication by and for dystonia investigators.

The scientific journal is the only open-access journal dedicated to dystonia research, serving as the central repository for all dystonia research.

In 2023, *Dystonia* published a special issue, *Models, Mechanisms, and Maturation in Developmental Dystonia*, which offers the most recent perspectives in the genetics, molecular, and network mechanisms underlying developmental dystonia. *Dystonia* articles published this year covered a wide range of dystonia types and research fields. Articles with the most views this year covered writer's cramp, tardive dystonia, and non-focal dystonia.

Dystonia is a resource for researchers, clinicians and patients and is partially supported by the Joan Miller Young Investigator Fund.

Access *Dystonia* and all published research articles from 2023 at:
frontierspartnerships.org/journals/dystonia

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DMRF Grants and Fellowships Further Dystonia Understanding and Treatment

*Committed to advancing dystonia research, DMRF supported **14 research grants, 5 research fellowships and 2 clinical fellowships** in 2023. The DMRF has invested more than **\$38 million in grants and fellowships** since 1976 in an effort to provide investigators with the funding needed to address the most pressing, unresolved questions in dystonia research.*

GRANTS

The research grant program supports the advancement of our understanding of the underlying mechanisms of dystonia; supports the exploration of new areas for drug discovery and allows investigators to pursue new technologies and how these can be applied to improving the lives of those living with dystonia.

GABA Abnormalities and Stability in Cervical Dystonia

Brian Berman, MD, Virginia Commonwealth University

Normalizing DYT1 Cholinergic Neurons by CRISPR Disruption of Mutant TOR1A Allele

Xandra O. Breakefield, PhD, Massachusetts General Hospital

Neuroanatomical Isolation of Networks in Dystonia Through Analysis of Causal Brain Lesions

Daniel Corp, PhD, Deakin University

Octanol Effect on Symptoms and Network Dysfunction in Myoclonus Dystonia: An Intoxication Free Therapy?

Cecile Gallea, PhD, Pitié-Salpêtrière Hospital

Personalized Functional Neuromodulation of Common Deficits in Focal Dystonias

Leighton Hinkley, PhD, University of California San Francisco

**A Next Generation Sensing Neural Interface Study
for Adaptive DBS in Dystonia**

Simon Little, MD, University of California San Francisco

Protein Kinase R Dysfunction in Dystonia

Stephanie Moon, PhD, University of Michigan

**Theta Burst Transcranial Focused Ultrasound
as a Novel Treatment for Cervical Dystonia**

Jean-Francois Nankoo, PhD, University Health Network

**Investigating Multimodal Neuroimaging for Probing Brain
Networks in Cervical Dystonia**

Richard Reilly, PhD, Trinity College Dublin

**Developing High-Throughput Assays to Enable Drug
Development for DYT1 Dystonia**

Christian Schlieker, PhD, Yale University

**Cerebellar Repetitive Transcranial Magnetic Stimulation
in Monogenic Myoclonus-Dystonia**

Anne Weissbach, PhD, University of Lubeck

Deciphering the Sources of Myoclonus in DYT-SGCE Patients

Yulia Worbe, MD, PhD, Pitié-Salpêtrière Hospital

Molecular Mechanisms of Dysmyelination in Dystonia

Dan Yellajoshyula, PhD, Case Western Reserve University

Mapping Anxiety in Cervical Dystonia

Abhi Mahajan, MD, Rush University

CLINICAL FELLOWSHIPS

The DMRF supports the training of young physician scientists—both to help them become better dystonia clinicians and to support their important clinical research. The DMRF has supported 34 clinical fellows since the program launched in 2012.

John Yu, MD

Mentor: Michael Okun, MD
University of Florida

Mariel Pullman, MD

Mentor: Rachel Saunders-Pullman, MD
Icahn School of Medicine at Mt. Sinai

POSTDOCTORAL RESEARCH FELLOWSHIPS

Postdoctoral fellowships support outstanding young scientists who have earned a doctoral degree and embarked on a period of mentored research. DMRF supports postdoctoral fellows who are working to fundamentally improve our understanding of brain dysfunction and molecular mechanisms underlying dystonia.

Investigating Abnormal Neurodevelopment in a Novel In Vivo Model of Inherited Dystonia

Simon Lowe, PhD, University College London Institute of Neurology

The Role of Basal Ganglia Pathways Engaged in Skilled Forelimb Movement in a Mouse Model of DYT1 Dystonia

Filipa França de Barros, PhD

Fundação D. Anna de Sommer Champalimaud e Dr. Carlos Montez Champalimaud

Beyond Theta: Analyzing Oscillations Across the Frequency Spectrum in Patients With Dystonia Implanted With Sensing-Enabled Pulse Generators

Stephanie Cernera, PhD, University of California San Francisco

Investigating Sensory Dysfunction in a Mouse Model of Early-Onset, Generalized Dystonia

Amanda Pocratsky, PhD, University College London

Mahlon DeLong Young Investigator Award

Spike-Triggered Adaptive Closed-Loop Cerebellar Deep Brain Stimulation (DBS) for Dystonia

Linda Kim, PhD, Baylor College of Medicine

DYSTONIA COALITION PILOT PROJECT PROGRAM AWARD

The goal of the Pilot Project Program is to foster the most promising clinical and translational studies of direct relevance to dystonia. Working collaboratively to support more research, the DMRF co-funded this grant with the Dystonia Coalition.

Genetic Characterization of GBA Mutations in Dystonia Using Long-Read Sequencing

Sebastian Schreglmann, MD, PhD

University Hospital Würzburg

“It is so important to build connections and friendships with others who understand what you are going through as someone with dystonia. DMRF support groups help provide those connections and offer practical help and experiences about living with dystonia that you typically can’t get from doctor visits.”

*Paula Schneider,
DMRF Vice President
of Support*

Support Groups Make Connections and Grow Friendships

DMRF support groups play a critical role in battling the isolation that dystonia often brings. By providing local peer support, support groups connect the community with others who understand the challenges of living with dystonia.



**DMRF has
48 support groups
and hosted 137
support group
meetings
in 2023.**

In 2023, DMRF hosted 137 support group meetings (94 virtual, 36 in-person and 7 hybrid.) The support network has grown significantly as individuals reach out to express interest in starting support groups in their area.

Last year, DMRF welcomed seven new groups to the support network, including Florida, Kentucky, Maine, Utah, New York (Rochester), Ohio (Columbus area), and Seattle.

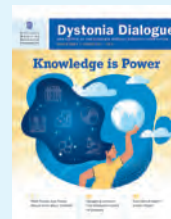
Support leaders are instrumental in fostering safe environments where people can share experiences, exchange information, and establish a support system they can turn to when in need. To encourage and educate support leaders, DMRF hosted two virtual support leader gatherings where leaders tapped into the knowledge and experiences of other support leaders from around the country.

Along with local support groups, DMRF’s moderated online forums, accessible through Facebook, provide dystonia support and information around the clock to people all over the world. Various DMRF online forum groups are available for specific areas of interest, such as dystonia and deep brain stimulation, parenting with dystonia, and dystonia spouses and loved ones. The largest online forum is DMRF’s Cervical Dystonia group with more than 12,000 members.



Dystonia Dialogue Informs and Connects

DMRF produced three issues of the *Dystonia Dialogue* magazine, which features updates on the latest dystonia research, personal profiles, coping tips for living with dystonia, and community news and events. With a circulation of 55,000, the *Dialogue* continues to be an integral platform for connecting the dystonia community.



Dystonia Zoo Days Expand to New Cities

The Dystonia Zoo Day events were a tremendous success in 2023 bringing together members of the dystonia community from around the country to raise awareness and funds for research. With a total of 11 in-person events and one virtual event, this year's Zoo Days saw expanded outreach.

**DMRF hosted
12 Dystonia Zoo
Day events in 2023,
including
5 new cities.**

Notably, five of the Zoo Days were hosted for the first time in Lansing, MI; Bridgeport, CT; Fort Wayne, IN; Indianapolis, IN, and Long Island, NY. The DMRF is exceptionally grateful to all the volunteers who worked tirelessly to organize these events.

DMRF hosted its 4th annual Virtual Zoo Day, providing the opportunity for people from coast to coast to come together on the same day at the same time to support one another and create awareness. The program included a panel of dystonia caregivers, mental health strategies, a discussion on the benefits of service animals, a zookeeper chat, and more.





DMRF Events Build Community

The DMRF community came together in so many fun, creative ways to promote dystonia awareness and raise funds for the Foundation in 2023. Along with Dystonia Zoo Days, DMRF network events raised more than \$500,000 in support of our mission to advance dystonia research, support those living with dystonia, and ultimately find a cure.



Dystonia Pet Palooza brought animal lovers together to raise awareness while celebrating the special role pets have in the lives of those affected by dystonia. Participants featured their pets and invited friends and family to make a gift in honor of their beloved pet.

DMRF Board members and father-daughter team, Allison London and Ron Hersh and their extended family hosted the inaugural **Positive Altitude Run/Walk** at the Central Park Reservoir in December.

#Riding4Us DMRF Board Member Jon Davis cycled 4,000 miles and completed a 6-day bike journey across Scotland to raise awareness about dystonia and funds for research.



Other network events included **Nashua Dystance4Dystonia 5K Walk/Run**, **SB Squares4Dystonia**, **Party with a Purpose**, **Dystonia Warrior Ride**, and the **Dystonia Tailgate**. Unfortunately, the **1st Angels Walk for Dystonia** in New Jersey was canceled due to a non-tropical coastal storm. Despite the setback, event organizers raised significant funds, spread dystonia awareness to their community, and have committed to another walkathon in 2024. DMRF extends a huge THANK YOU to all the volunteers who organized, fundraised, and participated in these outstanding events.



Dystonia Awareness Month Rocks Blue

The dystonia community rallied in 2023 during National Dystonia Awareness Month in September with two new awareness campaigns.

Shine a Light on Dystonia. The community embraced the idea of lighting up landmarks and buildings in blue for dystonia awareness by illuminating Niagara Falls; Chicago's Willis Tower; the Peace Bridge and Market Square's Spires in Rapid City, SD; Massachusetts Bridges, including Zakim, Longfellow, Fore River, and Burns Bridge; the San Diego Convention Center; and City Hall Houston.

Wear Blue in September. This campaign encouraged supporters to pick a day in September, or even one day each week, to don blue attire and wear stickers that said, "Ask me why I am wearing blue." Friends, family, co-workers, and corporate supporters participated in this visual show of support.

Niagara Falls



Baltimore



Minneapolis



Tackling Non-motor Symptoms of Dystonia

The DMRF has long recognized the need to better understand the non-motor symptoms of dystonia that can profoundly impact the lives of those affected. These hidden dimensions, including anxiety, depression, sleep disruption, chronic pain, fatigue, and cognitive impairment, can significantly diminish quality of life.

In an effort to focus attention on this important area, learn more, and encourage collaboration among the scientific community, the DMRF hosted a virtual scientific workshop, *Non-Motor Symptoms of Dystonia*, in March 2023. The meeting brought together experts from across the world to discuss the evidence of non-motor symptoms in dystonia, examine existing assessment and treatment methods, and to collaborate on future progress in the field.

The program was co-chaired by Kathryn Peall, MD, PhD of Cardiff University in the United Kingdom and Davide Martino, MD, PhD of the University of Calgary in Canada. Drs. Peall and Martino were so encouraged by the discussion at the workshop that they summarized the *Non-Motor Symptoms of Dystonia* workshop for the DMRF's journal, *Dystonia*. They also approached the International Parkinson

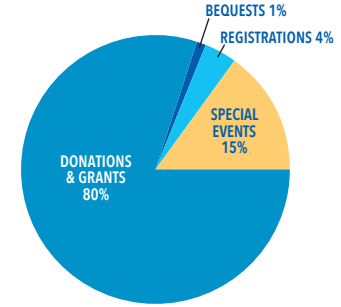
and Movement Disorder Society (MDS) about the importance of the topic, and it is now a study section with MDS.

Participants continue to meet on the topic, including an in-person meeting the DMRF arranged during the International Dystonia Symposium in Dublin last summer. Through their conversations throughout 2023, investigators reached consensus on a screening tool that can be used in the clinic to help physicians identify patients who are struggling with non-motor issues. This screening tool is currently being tested in four clinics in the UK and in North America with the plan to roll out testing to a wider group of centers after the feasibility stage. If approved, the screening tool will be a huge help to doctors and patients in addressing the non-motor symptoms that can be associated with dystonia.

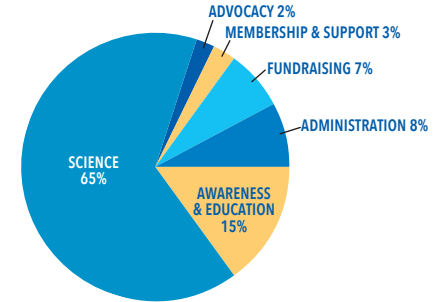
STATEMENT OF ACTIVITIES

Dystonia Medical Research Foundation | Year Ending December 31, 2023

REVENUES, GAINS & SUPPORT	WITHOUT DONOR RESTRICTIONS	WITH DONOR RESTRICTIONS	TOTAL
Donations and grants	\$ 2,009,404	\$ 262,760	\$ 2,272,164
Registrations	128,610	-	128,610
Cure Dystonia Now funds received	78,446	-	78,446
Bequests	34,937	-	34,937
Special events, net of direct benefits to donors of \$75,899	452,324	-	452,324
Investment income	493,677	-	493,677
Other income	3,776	-	3,776
Net assets released from restrictions	363,731	(363,731)	-
Total Revenue, Gains and Support	3,564,905	(100,971)	3,463,934
EXPENSES			
Program services -			
Science	2,215,520	-	2,215,520
Awareness and education	506,313	-	506,313
Membership and support	101,745	-	101,745
Advocacy	71,903	-	71,903
Total program services	2,895,481	-	2,895,481
Supporting services -			
Administration	286,313	-	286,313
Fundraising	225,674	-	225,674
Total Expenses	3,407,468	-	3,407,468
Change in Net Assets	157,437	(100,971)	56,466
Net Assets, Beginning of Year	2,642,291	2,153,862	4,796,153
Net Assets, End of Year	\$ 2,799,728	\$ 2,052,891	\$ 4,852,619



REVENUES



EXPENSES

STATEMENT OF FINANCIAL POSITION

Dystonia Medical Research Foundation | Year Ending December 31, 2023

ASSETS	2023
CURRENT ASSETS	
Cash and cash equivalents including segregated	
Dystonia Coalition cash of \$601,276	\$ 689,422
Unconditional promises to give	529,047
Other current assets	24,573
Finance lease right-of-use asset, current	4,819
Operating lease right-of-use asset, current	42,681
Total Current Assets	1,290,542
 NONCURRENT ASSETS	
Investments	4,975,677
Equipment and leasehold improvements,	
less \$35,194 of accumulated depreciation	8,263
Finance lease right-of-use asset, noncurrent	3,756
Operating lease right-of-use asset, noncurrent	165,456
Total Noncurrent Assets	5,153,152
Total Assets	\$ 6,443,694

LIABILITIES & NET ASSETS	2023
CURRENT LIABILITIES	
Grants and fellowships payable	\$ 563,470
Accounts payable and accrued expenses	93,240
Deferred revenue	75,000
Dystonia Coalition agency liability funds	601,276
Financing lease liability, current	4,818
Operating lease liability, current	49,299
Total Current Liabilities	1,387,103

NONCURRENT LIABILITIES	
Financing lease liability, noncurrent	3,756
Operating lease liability, noncurrent	200,216
Total Noncurrent Liabilities	203,972
Total Liabilities	\$ 1,591,075

NET ASSETS WITHOUT DONOR RESTRICTIONS	
Undesignated	1,299,728
Board designated for science	1,500,000
Total net assets without donor restrictions	2,799,728

NET ASSETS WITH DONOR RESTRICTIONS	
Purpose restricted	2,052,891
Total net assets with donor restrictions	2,052,891
Total Net Assets	4,852,619

Total Liabilities and Net Assets	\$ 6,443,694
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*A complete copy of financial statements audited by
 Barnes, Givens & Barnes, Ltd. is available upon request from
 the Dystonia Medical Research Foundation,
 One E. Wacker Drive, Suite 1730, Chicago, Illinois 60601.*

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\$25,000 OR MORE

Team AbbVie
Team Ipsen Biopharmaceuticals, Inc.
Team Merz Therapeutics

\$5,000 TO \$9,999

Team Kristin/Toledo
Team Revance Therapeutics Inc.

\$2,500 TO \$4,999

Team Casler/Lansing
Team Erinn's Explorers/Portland
Team Fight for a Cure/Long Island
Team Hinsdale Bank and Trust/Chicago
Team Not Josting Around/Boston
Team Metronet/Lansing
Team TWISTED BADDIES/Virtual

\$1,000 TO \$2,499

Team A Twist of Ed/Pittsburgh
Team Allegheny Health Network (AHN)/Pittsburgh
Team Aidan/Indianapolis
Team Alta Equipment Group/Toledo
Team Braden's Brigade/Pittsburgh
Team Carson Strong/Ft. Wayne
Team Eisel/Toledo
Team Hartford Healthcare/Connecticut
Team Henry Ford Movement Disorders/Toledo

Team Impact Networking/Chicago
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The background features a light blue color palette with a pattern of hexagons. Some hexagons are solid, while others are outlined. Overlaid on this pattern are several wavy, multi-line patterns in various shades of blue, creating a sense of movement and depth. The overall aesthetic is clean, modern, and scientific.

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