HIGHLIGHTS FROM 2022

CONTINUED OUR DISSECTING DEVICES WEBINAR SERIES
This free educational program profiled six products in the epilepsy technology space. Over two years, this information has reached more than 1,000 patients and caregivers.

13TH ANNUAL EVENING OF HEARTS & HUGS EVENT
Our annual gala achieved record-setting fundraising numbers. Staged from Theater on the Lake, supporters participate both in Chicago and by watching virtually from across the United States.

CO-HOSTED 3RD ANNUAL DEVICE ACCELERATOR COURSE
In collaboration with the Epilepsy Foundation, we facilitated a six-week virtual course to provide resources and experts for five new epilepsy device companies.

DIVERSITY RADIO CAMPAIGNS CHICAGO, DETROIT AND DC

PARTNERS AGAINST MORTALITY IN EPILEPSY (PAME) CONFERENCE
Danny Did co-hosted a SOLD OUT meeting that was attended by more than 250 bereaved caregivers, healthcare professionals, researchers, and advocates to advance progress against mortality in epilepsy.

CHARITY PARTNER TO THE HIGHLAND PARK HIGH SCHOOL CHARITY DRIVE
A highlight of our spring was our partnership with the Highland Park High School Charity Drive. More than 2,000 students and staff spent 5 weeks raising epilepsy and SUDEP awareness throughout the school and Highland Park, IL community. This effort raised nearly $100,000.

IN-STORE AWARENESS & FUNDRAISING CAMPAIGN IN OVER 300 WALGREENS LOCATIONS

Partnering with Walgreens to Protect Kids with Epilepsy
Your $1, $3 or $5 donation makes a big difference. Give to the Danny Did Epilepsy Foundation at your Walgreens Counter.
# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>Page</th>
<th>Section</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Meet Danny Stanton</td>
</tr>
<tr>
<td>7</td>
<td>Letter From the Co-founders</td>
</tr>
<tr>
<td>8</td>
<td>Letter From the DDF President</td>
</tr>
<tr>
<td>9</td>
<td>Mission, Vision &amp; Values</td>
</tr>
<tr>
<td>10</td>
<td>About Epilepsy &amp; SUDEP</td>
</tr>
<tr>
<td>12</td>
<td>2022 Year in Review</td>
</tr>
<tr>
<td>22</td>
<td>Our People</td>
</tr>
<tr>
<td>29</td>
<td>Family Ambassador Program</td>
</tr>
<tr>
<td>31</td>
<td>Get Involved</td>
</tr>
</tbody>
</table>
MEET DANNY

Danny Did Foundation
Danny Stanton was born on March 2, 2005. Danny died on December 12, 2009.

A wonderfully normal two years of development followed his birth as Danny joined his sister Mary Grace and his brother Johnny and took his place as kid number three in the Stanton family. He was, as his grandma Murtaugh dubbed him, the “little toughie” of the family. The kid was a scrapper who held his own as he established his place within his family. Even at two years of age, he had established himself—somehow, and in many ways—as an equal not only in terms of experiencing life, but also as one who brought experience to the lives of everyone in his family. Soon, Danny was no longer the youngest Stanton, as his brother Tommy came along when Danny was two.

It was at age two when Danny had his first seizure. His eyes rolled back and he shook. We did not know what was happening. We dialed 911 then rushed him to our next-door neighbor, who is a paramedic. Danny seized for almost 10 minutes. On that night, we had a battery of tests done at Children’s Memorial Hospital in Chicago. There was no known reason or cause found for Danny’s seizure. No injury, no trigger. We were told that sometimes seizures in children have no known cause and that kids often outgrow them. The experience was terrifying.

We left the hospital that night without any written information about seizures or seizure disorders, sans any mention of epilepsy or epilepsy organizations, and sans anything short of the feeling that we had fallen into a hole that had no bottom. Sudden Unexpected Death In Epilepsy, of course, crossed no lips and fell upon no ears. Danny began sleeping in bed with us after that first seizure.

Danny had a second seizure while sleeping just over a month later. After this second seizure, he had an MRI. Again, no underlying cause was found for his seizures. Danny was then put on a variety of medicines to control the seizures, and regular electroencephalograms (EEGs) were scheduled and performed. His neurologist determined that Danny had childhood seizure disorder. Still no mention of epilepsy, still no suggestion of outreach to epilepsy organizations, and still, of course, no mention of Sudden Unexpected Death In Epilepsy. What we were told was this: he’ll outgrow it, many kids experience this, we don’t know why it happens, maybe it’s “his normal.” How might life be different had a sophisticated and tough-minded doctor sat us down and said “let me tell you something that will shock you but that I am mandated both by my profession and by my personal sense of duty to inform you of . . . your son has epilepsy, and along with this diagnosis comes a plethora of safety issues that you need to become aware of…”

Danny’s daily behavior was completely normal. He continued to be athletic, inquisitive, engaging, social, and, more than anything, affectionate. As we continued meeting with Danny’s neurologist for regular check-ups, we became more and more comfortable with the idea that we were doing all we could for Danny and that we were on a clear and well-defined path. We were told that a slightly abnormal EEG might just be Danny’s ‘normal.’

After sleeping with us for a half a year, Danny began to return to his own bed. On hundreds of occasions we left our bed to check on him. We caught him twice more having a seizure in his sleep. We informed our doctors of every event.
We continued to trust and believe that we were doing all that we could. Birthdays, holidays, block parties . . . each month brought more punch to us as a family, and each day brought hugs from Danny: he’d tuck his arms into his own chest and then wait for you to surround his body with yours. He’d then nestle his head under your chin, take a deep breath, and let his exhale say that he had found the spot.

After a year of no known seizures, Danny returned again to his own bed. And then as he turned four and had gone without a known seizure for 18 months, we thought that our course of action had been successful and that his neurologist was right: Danny would, through medicine and time, outgrow his ‘seizure disorder.’ It was like a rebirth for us and for Danny.

On Tuesday, December 8, 2009, we took Danny for a scheduled appointment with his neurologist. He had a one-hour EEG done while sleeping, and we met with his doctor afterward. There was still abnormal activity showing up, but maybe a lesser amount than the previous EEG showed. We increased Danny’s dosage that day to accommodate a slight weight gain since his last visit.

Four days later, on December 12, 2009, we found Danny’s lifeless body in his bed at 7:15 am.

How many seizures did Danny have that we did not catch? We will never know. One thing we do know are the last words to ever fall upon that little toughie’s ears. They are the words whispered by his mom to Danny each and every night as she put him to bed: “I love you, sweet boy.” These words were always followed by three kisses: one to the forehead, one to the nose, and one to the lips. I love you, sweet boy.

Now life continues –in many respects– albeit without the soft bounce of one little boy’s feet upon the dirt of the Earth. Our Danny boy has entered a new phase, part of which is in the form of his organization, the Danny Did Foundation, which derives from the last line of Danny’s obituary: Please go and enjoy your life. Danny did.
Im imperfectly, our son Danny had epilepsy. Tragically, our son Danny died from epilepsy. Specifically, Danny died from Sudden Unexpected Death in Epilepsy, also known as SUDEP. Graphically, Mariann found Danny dead in his bed one cold December morning two weeks before Christmas in 2009. Horrifically, we tried to resuscitate Danny that morning in front of his sister Mary Grace and his two brothers, Johnny and Tommy. Distressingly, we believed that we had been doing everything possible to care for Danny.

Through two years of treatment by medical professionals, we were never informed that Danny had epilepsy, and even though Danny suffered from seizures we were never informed that Danny could die from one of his seizures. We were never told about SUDEP. We were never given information about organizations that could educate and support us as we swirled through Danny’s seizure episodes.

And so when Danny died, we knew that he and that we had been robbed of the opportunities of a lifetime. An actual lifetime. And we were furious. And we asked each other in our bedroom on that morning in December – after we had returned home from the hospital were Danny’s body had grown cold atop of ours as we laid with him - we asked ourselves, “how could this happen?”

Part of our answer to that question has been the formation of the Danny Did Foundation, which we are nurturing and developing as if it were part of Danny himself, and together with so many other individuals and organizations, we are changing the perception of epilepsy and the outlook for those afflicted by epilepsy.

And so as one year closes and another begins, we look forward to new partnerships and to new alliances as we continue our work to enable access to seizure monitoring and detecting devices for families who cannot afford them, and to advance awareness of epilepsy and the risks that accompany it, including Sudden Unexpected Death in Epilepsy. Thank you for your participation and for helping to prevent another death caused by a seizure.

Mariann and Mike Stanton
Letter from the DDF President

The year 2022 was full of progress and excitement at Danny Did. We kicked off with a new partnership with the Highland Park High School Charity Drive. More than 2,000 students learned about epilepsy and became energized ambassadors for our work. They also mobilized to raise meaningful funds to expand our reach to more families who face epilepsy. We also began a new relationship with one of the fastest-growing companies in Chicago - Redwood Logistics - who empowered their entire global workforce to raise awareness and funds for Danny Did as their official 2022 charity partner.

This year, my nephew Danny would have been a senior in high school, experiencing a time of excitement when his curiosity, talents and friendships would have grown with the passing of another school year. Epilepsy impacts 1 in 26 Americans, and is second only to stroke among neurological disorders in Years of Potential Life Lost (YPLL). Each family who has lost a loved one to epilepsy knows how painful those lost years can be. The communication between medical professionals and their patients about risk of SUDEP, while improving, is still unacceptably low.

Over this summer, Danny Did had the privilege to present during the International League Against Epilepsy’s European Epilepsy Congress session on "How To Talk About SUDEP." This was a big opportunity to emphasize to providers across Europe the importance of having open discussions about the risks of epilepsy mortality. This partnership included Lauren Knepper, the Co-President of our DDF Associate Board, who shared her perspective as an adult patient; and Dr. Stephan Schuele, head of the Northwestern Memorial Hospital Comprehensive Epilepsy Center, who encouraged his peers to talk about SUDEP. Danny did reach Europe this year.

As we look back at the past 12 months, what makes us most proud is that our support for families in need has extended to all 50 states and 15 countries, support that many have described as life altering. As we grow to help more people, we encourage and appreciate your partnership. We are excited and inspired to drive more progress.

With hope and gratitude,

Tom Stanton
To advance awareness of epilepsy and Sudden Unexpected Death in Epilepsy (SUDEP), protect people with epilepsy, and prevent deaths caused by seizures.

To create a reality in which healthcare providers openly communicate SUDEP and all mortality-related risks with their patients or caregivers.

To elevate awareness of, and access to, technologies that provide early intervention and added safety when a seizure occurs.

Mission

Vision

Values

Family Love: Everything we do is in service of families;
Preparation: We help families prepare to protect their loved ones;
Hope: We hold hope for a brighter future;
Enjoyment of Life: Our work is meant to carry forward and spread Danny's joy for life.
A seizure is a sudden behavioral change resulting from excessive electrical activity in the brain. Seizures may be provoked (also known as acute symptomatic; for example, a fever in a young child, drug intoxication or withdrawal, electrolyte imbalance) or unprovoked, resulting from a genetic, developmental, acquired (for example, head trauma, tumor, stroke), or an unknown cause.

When a person has two or more unprovoked seizures, they have epilepsy. An estimated 3.4 million Americans have active epilepsy, with up to 200,000 new cases diagnosed in the United States each year. Approximately 1 in 26 people will develop epilepsy at some point in their lives, and the onset of epilepsy is highest in children and older adults. Epilepsy impacts 65 million people worldwide and is a spectrum that spans more than 40 syndromes. Despite all available treatments, 30 to 40 percent of people with epilepsy continue to experience uncontrolled seizures. In two-thirds of patients diagnosed with epilepsy, the cause is unknown.

A report from the Centers for Disease Control and Prevention (CDC) indicates that the number of Americans with epilepsy is higher than ever. Epilepsy affects more people than multiple sclerosis, cerebral palsy, muscular dystrophy, and Parkinson’s combined – yet receives fewer federal dollars per patient than each of these. According to advocate group Citizens United for Research in Epilepsy, epilepsy costs the United States approximately $15.5 billion each year. The indirect costs associated with uncontrolled seizures are seven times higher than that of the average for all chronic diseases.
Seizures can sometimes be fatal from a range of causes, and no matter how the death is defined, every time it is a huge loss. There are numerous ways that a person can die stemming from seizure activity – drowning, an accident, head trauma, suicide, or status epilepticus. Another category of epilepsy mortality is Sudden Unexpected Death in Epilepsy, or SUDEP. SUDEP is the most common category of death in epilepsy. SUDEP is just that: sudden and unexpected. The most recent research shows that a case of SUDEP occurs in 1 in 1,000 people with controlled epilepsy, and for 1 out of 150 people with uncontrolled epilepsy.

The Danny Did Foundation pledges to shine a light on SUDEP for as long as is necessary to achieve major progress in awareness and in the advancement of preventative measures.

WHAT DEFINES A CASE OF SUDEP?

SUDEP is not a cause of death but rather a category of death. A death is referred to as SUDEP when a seemingly healthy person with epilepsy dies unexpectedly and no clear reason for the death can be determined. In most cases, an autopsy is required to rule out other causes of death. The most common criteria used to determine whether a death is due to SUDEP are:

- The person has epilepsy, which is defined as recurrent unprovoked seizures.
- The person died unexpectedly while in a reasonable state of health.
- The death occurred suddenly and during normal activity (often during sleep).
- An obvious medical cause of death could not be determined at autopsy.
- The death was not the direct result of status epilepticus.
2022 YEAR IN REVIEW

Danny Did Foundation
Awareness Campaigns

13 Campaigns in 2022

For a collective reach of more than 8.5 million people!
Our practical impact is to provide seizure alert devices for families in need. These systems provide added safety for patients, and enhanced peace of mind for caregivers.
STORIES OF IMPACT
Martinez Family, Texas

"I can't thank you guys enough for the grant that has helped us sleep better at night, knowing that if Kimberly has a seizure we will be notified. This is something a parent is always worry about, thankfully we heard about your foundation and with your help we were able to get the Sami camera. My husband and I used to take turns sleeping in the same room as Kimberly, just to make sure she was safe. My family is very thankful for The Danny Did Foundation, you guys are doing God's work helping ease the worry out of parents with kids that have epilepsy."

Newton Family, Canada

"I wanted to thank you and the foundation again for making this possible for my family. Piper was sick this past weekend with a high fever, and I’ve never slept so well knowing that the SAMi would alert me if anything happened! I was so incredibly thankful to have the monitor. So thank you again for everything you do!"

Dailey Family, Texas

"My son received the SAMi and it’s been a blessing to me because he sleeps so much better. And so do I, knowing we have extra help in detecting his seizures. I thank God for The Danny Did Foundation."
I am a doctor who attended the SUDEP symposium that Danny Did Foundation was a partner to with the Child Neurology Foundation. Listening to the speakers there completely changed the way that I talk with my patients about the risk of sudden death, and I now understand the importance of having the conversation with EVERY family. Since adjusting my communication on this topic, I have noticed an unexpected benefit: my patients are more compliant with their medications and more open about breakthrough seizures. I wish I had thought of this sooner! I have been using the materials provided at the symposium, and I will be ordering more soon.

For its work on SUDEP education and epilepsy care, the Danny Did Foundation is an outstanding organization and a Chicago jewel!
Educational
33 Presentations in 2022

- Rare Epilepsies Conference
- Chicago Epilepsy Centers Meetings
- Medical Students Attending the University of Illinois
- ILAE European Epilepsy Conference
**DO IT FOR DANNY FITNESS CLASS**
We took the action outside with a bootcamp fitness class in La Grange, Illinois. Led by FitGive, more than 100 participants worked up a sweat for kids who face epilepsy.

**HIGHLAND PARK CHARITY DRIVE**
More than 2,000 students and staff spent 5 weeks raising epilepsy and SUDEP awareness throughout the school and Highland Park, IL community. In addition education spread, this effort raised nearly $100,000.

**LOYOLA UNIVERSITY WOMENS VOLLEYBALL AWARENESS GAME**

**13TH ANNUAL EVENING OF HEARTS & HUGS**
Our record setting Hearts & Hugs gala welcomed guests to Chicago’s lakefront both in person and virtually from around the U.S.

**2ND ANNUAL PUSH UP CHALLENGE GOLF OUTING**
Lead by Co-Captains Kevin Cohen and Nathan Wysocki, this sold out event raised nearly $200,000 in Hawthorn Woods, IL.

**CHARITY PARTNER OF REDWOOD LOGISTICS**
TEAM DANNY DID IN 2022

AUGUST - SEPTEMBER
DIANE GARVEY SWIM MARATHON

OCTOBER
TEAM DANNY DID IN THE CHICAGO MARATHON

AUGUST
MIKE LITTAU CHICAGO TRIATHLON

SEPTEMBER
ANNUAL PUSH UP CHALLENGE

DECEMBER
SETH GRANDA IN HUNTSVILLE, AL MARATHON
MEDIA COVERAGE

Loyola University Volleyball Player Shares Story to Advocate

17-Year-Old Earns Master Chess Title After Four Brain Surgeries

Advancing Technology Solutions in Pediatric Epilepsy

Walgreens In Store DDF Campaign Raises Funds to Fight Epilepsy

Understanding Sudden Expected Death in Epilepsy

DDF Ambassador Raises SUDEP Awareness for Late Daughter
OUR

PEOPLE

Danny Did
FOUNDATION
Mike grew up in a family of 10 in Chicago’s Rogers Park neighborhood. After graduating from Loyola Academy, he earned his college degree from Northwestern University. He has worked in law enforcement for more than 20 years. He and his family live on the northwest side of Chicago.

Mariann grew up near Chicago’s south side in Oak Lawn, Illinois. Mariann attended Mother McCauley high school, where she was a star basketball player before going on to Loyola University as a four-year scholarship basketball player. She earned a degree in English from Loyola and then went on to earn a Master’s degree in Education from DePaul University. Mariann is a high school teacher in Chicago.
Tom became Danny Did's founding executive director in 2010. He remembers many endless summer days and nights playing with Danny and his siblings on Leona Avenue. Tom leads the communications and fundraising efforts for Danny Did, and works to expand their collaborations and impact in the larger epilepsy community. Tom holds a journalism degree from the University of Dayton, and a master’s degree in Public Policy from DePaul University. More than anything, Tom gained happiness from the love Danny gave by way of lots of hugs. He is thankful to have the chance to keep Danny’s spirit alive through service to Danny Did.

Tom Stanton  
President

Mary has been a part of the DDF staff since 2010. However, her relationship with Danny’s family began when her young sons, Brenden and Charlie, spent years playing sports with the Stanton children. Mary leads Danny Did’s efforts with family support, event planning, and volunteer coordination. She earned her Bachelor’s of Science at Northern Illinois University and her Master’s in Human Resources at Loyola University in Chicago. Mary considers it an honor to serve atDanny Did, and the Stanton family is most grateful not only for her friendship, but also for the wonderful buddy that Charlie was to Danny.

Mary Duffy  
Executive Director

As the Danny Did Programs Coordinator, Katie has the role to make a direct impact on families who reach out in need. Her career has included time as a paramedic, where she gained experience in treating those who had suffered a seizure. Prior to working at Danny Did, Katie was a long-time volunteer for the organization, and she serves on its Associate Board. As a former neighbor to the Stantons, she held a special friendship with Danny.

Katie Gaughan  
Programs Coordinator
Julie is a media sales executive and her career path has featured sustained success at some Chicago’s largest radio stations. She is currently an executive at Prime Time Media, where she utilizes her experience and relationships to connect the Danny Did Foundation with a variety of awareness platforms and opportunities. Julie is a graduate of Purdue University, and lives in Wilmette, IL with her husband and their children.

Lynn is an attorney specializing in commercial real estate development, land use and zoning. Lynn has also taught real estate and zoning classes at the University of Richmond Law School and the Loyola University of Chicago Law School. She holds a BA from Miami University, an MA from Xavier University and a JD from Loyola University of Chicago. She and her family live in Winnetka, IL.

When the eldest daughter of Tracy and Brian Jablonski was diagnosed with epilepsy and a brain tumor in 2015, they learned about the Danny Did Foundation. She has since become an advocate to spread awareness about epilepsy, SUDEP, and the technologies available. Tracy is a graduate of St. Francis University. She and her family live in LaGrange, IL.

Dr. Garcia-Sosa is an Attending Physician in and the Epilepsy Center at Ann & Robert H. Lurie Children’s Hospital of Chicago, and Assistant Professor of Pediatrics (Neurology), Northwestern University Feinberg School of Medicine. She represents Danny Did for educational speaking engagements at with medical students, family events and medical conferences. She is a native of Puerto Rico.

Jerry is a physician at the University of Illinois College of Medicine at Chicago. He first started working with Danny Did as an executive board member of Northwestern University’s Dance Marathon, where he successfully managed an effort that raised $1,214,632. Jerry also holds a B.A. from Northwestern and an M.S. in from Georgetown. He is a native of Buffalo Grove, IL.
DOUG BRUNO - EMERITUS

Doug Bruno has more than 700 career victories as head coach of DePaul University’s Women’s basketball. In addition, as a coach for USA Basketball, he coached two different teams to Olympic Gold Medals. Doug was a founding board member for the Danny Did Foundation. He now serves as a member of our Emeritus Board. His family knows the challenges of epilepsy.

JOSH GABBY - TREASURER

Josh is a data analytics executive for JPMorgan Chase. His support for Danny Did includes strategy planning, marketing expertise and service as our DDF board treasurer. He holds an MBA from Northwestern University’s Kellogg School of Management.

DIANE GARVEY-SECRETARY

Diane has hosted several fundraising events for Danny Did, with several featuring her role as a Zumba dance instructor. This year she completed a 26.2-mile swim-a-thon in her North Carolina community to benefit kids who face epilepsy.

JOHN WRENN

John owns and operates several Chicago area businesses, including Theater on the Lake, a premier dining and event space located on Chicago’s lakefront in Lincoln Park. He is active in all forms of philanthropy and specializes in fundraising and development.

KARI HEIDEMAN

Kari is a family nurse practitioner who brings a valuable healthcare provider perspective to the Danny Did board. She has also served as Co-Chair of our annual Hearts & Hugs gala, and is a repeat member of Team Danny Did.

MIKE STANTON - CO-FOUNDER

With his wife Mariann, Mike is the co-founder of the Danny Did Foundation. His vision for keeping Danny’s spirit alive in the world through the work of the foundation is a guiding light. He is a repeat member of Team Danny Did. He does regular public speaking engagements on behalf of the foundation.

Danny Did Foundation
Margaret Storey is Professor at DePaul University and a parent advocate for children with medical complexity. She comes to epilepsy advocacy as the parent of daughter with Aicardi Syndrome, a rare and debilitating condition that causes early-onset, intractable seizures. She received her PhD from Emory University, and has written for The Guardian online, the Motherlode Blog at The New York Times, and The Chicago Tribune. She and her family live in Evanston, IL.

Caroline Mcateer - Vice President

Caroline has worked at DePaul University for over 20 years and is currently a Director of Development in the Office of Advancement. Caroline was diagnosed with epilepsy when she was a teenager. Her personal experience has led to her desire to raise awareness of epilepsy. She holds a BA from the University of Dayton and an MS from DePaul University. She and her family live in Chicago.

Jeanine MCShea

Jeanine has spent 25 years in the real estate industry, both as a broker and an executive. With her daughter Erin McShea, she oversees The McShea Group at @properties. Jeanine is mother to two adult children with epilepsy, both who serve on the Danny Did Young Professionals board. She holds a bachelor’s degree from Winona State University, and lives in Chicago with her family.

Jennifer Sime

Jennifer is a mother, wife and registered nurse in Decorah, IA. She advocates in honor of her son Brendan, who died from SUDEP in 2017 at age 19. Among her many forms of advocacy, she has run multiple marathons as a member of Team Danny Did.

Kurt Florian - President

Kurt is a lawyer and a past executive director for the Epilepsy Foundation of Greater Chicago. He advocates in honor of his adult daughter, and the many families he has supported over the years. He and his wife Linda live in Wheaton, IL.

Scot Gillespie

Scot is an Executive Vice President and GM in the Commerce Cloud division of Salesforce. He holds an undergraduate degree from National Louis University, and an MBA from Northwestern University’s Kellogg School of Management. Scot and his wife live in Woodstock, Illinois.
Our Danny Did Associate Board integrates young professionals into the varied programs and campaigns of the Danny Did Foundation. Through volunteer activities, social and fundraising events and networking opportunities, this Board encourages the next generation of leaders to engage in our mission to prevent deaths caused by seizures.

Kristin McShea - Co-President
Lauren Knepper - Co-President
Katie Onsager
Jen Stoner
Alexis Flores
Natalie Knepper
Mike Wojtychiw
Katie LaZarra Gaughan
Kaitlyn Schubert
Toyosi Ogunlana (Kansas City)
Anna Radoff
Erin Gard
Justin McShea (Grand Rapids, MI)
Katie Amys (Minneapolis)
Rose Broccoli

Jenni Florian
Liam Stanton
Scott Sandler
Max Paternoster
Jimmy Schmitz
Talia Shear
Ellie Rich
Belicia Espinal
Nazree Williams
Kiernan Dunham
Adriana Kitchens (Kansas City)
Page Syvertsen (Boston)
Katie Vogt (San Francisco)
Patrick Lazarra (Minneapolis)
Ross Gordon (New York City)
Diamanta Panford (Cleveland)
We Can't Do It Without You!

**SUPPORT DANNY DID**
1. Make a Tax Deductible donation at Dannydid.org
2. Utilize your company match program after you donate.
3. Purchase DDF gear at Dannydid.org
4. Donate auction Items

**ADVOCATE WITH DANNY DID**
1. Volunteer at a DDF Event
2. Ask your doctor to talk about SUDEP prevention and research
3. Share the Seizure Safety quiz at your School
4. Lead a “fan raising” Social media Campaign

**GET INVOLVED WITH DANNY DID EVENTS**
1. Organize and Host your own Fundraising Event
2. Sponsor a Danny Did Event
3. Coordinate a Seizure Safety presentation at Your School
4. Join Team Danny Did

**PARTNER WITH DANNY DID**
1. Medical Professionals: Help Us Talk About SUDEP
2. Families: Assist Danny Did’s Outreach to Doctors
3. Device Companies: Tell us about your technology

Find out more at Dannydid.org/get-involved
Thank you for your support!

Protecting Kids with Epilepsy

Danny Did
Foundation

@Danny_Did_Foundation

@dannydid.org

Please go and enjoy your life.
Danny did.