Our Mission

Jett Foundation empowers people and families impacted by Duchenne muscular dystrophy through the development of transformative programming, educational opportunities, and ongoing support for every stage of a Duchenne journey.

Letter from
the Executive Director

Jett Foundation spent this past year reconnecting on the ground with our Duchenne muscular dystrophy community... finally! Our team experienced so much joy and captured many special memories from those long-awaited, in-person reunions across our programs. From Camp Promise and Family Workshops to athletic events and fundraisers, connecting with our friends and supporters again was nothing short of amazing.

Our virtual programs have been critical to families over the past few difficult years. Moving forward, hybrid offerings of our programs and services will continue as our community comes together once again. Access to resources both online and in person ensures that the families we partner with across the nation benefit fully from our wrap-around services year-round.

As we have rebuilt and redefined how our programs are delivered long-term, several of the initiatives set forth in our Strategic Plan were accomplished in 2022. We launched a new website to better showcase our programs and be a living resource for our families and community members. We worked closely with adults in the community through Town Hall meetings to build out our programming for 18+ individuals. And, our dedicated professional team has grown in the past year. Our team continues to work tirelessly to identify support gaps and develop programming that meets the needs of families at every stage of their Duchenne journey.

This year, we received the exciting news that Jett Foundation is the recipient of a $1 million endowment gift from Karen and Rob Hale through the Fox Rock Foundation. The Hales’ support of our mission will ensure Jett Foundation’s deep-rooted impact in the community for years to come as we work toward long-term goals identified in our Strategic Plan.

Thank you for the role you have played in our organization to make a difference in Duchenne—we are so grateful for you. Whether you are a family utilizing our programs, athlete fundraising, donor, or volunteer cheering us on, your contribution to Jett Foundation has a meaningful impact in our community.

Sincerely,
Eric Snyder, Executive Director
Delivering Happy Holidays on Wheels: Two Families Receive Vans Just in Time for the Holiday Season

Andrew Simpson (15) lives with Duchenne muscular dystrophy and is cared for by his mother, Heather. They also live with Andrew’s great Uncle Jim and their three dogs, Moose, Pearl, and Daisy, in Washington State. The Simpson family enjoys engaging with their community and attending events that Andrew likes, but a lack of safe and accessible transportation made it challenging to do so. “It takes difficult transfer maneuvers to transition Andrew from a non-accessible car to his wheelchair and back,” says Heather. The family was planning their weeks far in advance in order to prioritize which appointments or events were worth the difficult transfers.

Heather and Andrew engage with Jett Foundation through awareness days and support groups regularly, but never imagined they would become participants of the Jett Giving Fund’s Accessible Vehicle Fund. “We watched families reach fundraising goals for their dream accessible vehicle and I never imagined that Andrew and I would be picked to be part of this life-changing program. Now that I look at our dream vehicle parked in the garage, I am so grateful that my friend pushed me to apply to the Accessible Vehicle Fund,” says Heather.

When Monday, November 13, 2022 arrived, Andrew and Heather could not contain their excitement for what was to come that evening. “My stomach was full of butterflies and excitement the entire day as I am sure Andrew’s was too! When I got home from work, Andrew was already ready for me to transfer him to the car. When we first saw the van, there were automatic tears in my eyes and Andrew let me know that the transfer would be the last time that I would need to lift him from the car to his chair. It still does not feel real.” Before receiving their accessible vehicle, Heather would transfer Andrew to their car, take apart his wheelchair, place it in the vehicle, and put it back together at their destination before lifting him back into the wheelchair and repeating the process when they left. “Now, Andrew is able to participate in after-school activities and we are starting to make appointments for things that he needs. We are so excited to get back into the community to go shopping at Target and local thrift shops,” says Heather.

Just a month later, on the opposite side of the country, the Fritz family shared a similar joy just in time for the holidays.

Lauren and John Fritz are parents to Ryan (16) and Jake (19) who live with Duchenne muscular dystrophy. The Pennsylvania family has experienced an increase in challenges, particularly in transporting their boys, as Jake and Ryan grow and become less ambulatory.

“We were attempting to lift and transfer Ryan, who is a full-sized young adult, to the back bench of our old van since it was only equipped for one wheelchair. There were several times that Ryan, my husband, or myself were injured in this process. It became impossible for us to travel safely together as a family.” - Lauren Fritz

On December 13, 2022, the Fritz family received their new vehicle that is both safe and accessible for Ryan and Jake’s wheelchairs, allowing the family to once again travel as a unit. To Lauren, the delivery of their van was a metaphor for life with Duchenne: both include challenges and patience, but there is an abundance of joy that comes with it.

“We all felt that Christmas had come early. It was wonderful to see the van pull into our driveway and know that we can once again experience a life of driving safely together.” - Lauren Fritz

For other families in the Duchenne community, Lauren shares: “There are many helpful people that have your back when you are attempting to secure something that is so fundamental to your family, such as an accessible vehicle. Our new van is not only a gain for the four of us, but for our entire community to be exposed to this resource. Jett Foundation and many others work hard to improve our lives and understand that what may seem small can actually make a major difference for your family.”
A Connecticut father reached out to Jett Foundation for assistance with putting food on the table for his family. Jason is a proud father of two boys. One of his sons, Justin (12), was diagnosed with Duchenne muscular dystrophy at just 18-months-old.

The Emergency Fund is an arm of Jett Foundation’s Jett Giving Fund, a financial support program for families impacted by Duchenne.

Jason first heard of the Emergency Fund program by attending Jett Foundation’s Patriots Training Day Camp with Justin a few years back. After Jason lost his job due to the COVID-19 pandemic and underwent other hardships, groceries became another costly expense on top of rent and medical bills.

“I barely can afford groceries, but I make sure my boys have food when they’re with me,” said Jason.

With help from the Emergency Fund, Jason and his family received aid for much needed essential items. To other families looking to apply for assistance, Jason says, “Keep up the fight. There are good organizations and people out there willing to help.”

“"The Emergency Fund grant has kept us going for a couple more months and helped us catch up on bills. It was a lifeline when we were drowning, and we will forever be grateful for the respite that Jett Foundation gave us.” - Rick Futrell

Jett Foundation approved Emergency Fund grants for these types of support in 2022:

- Food Assistance: 32%
- Rent or Mortgage: 22%
- Utility Bills: 34%
- Other: 13%

Established in 2017, Jett Giving Fund has helped transform the lives of countless families across the country with over $3.5 million worth of accessible equipment and emergency assistance thanks to the generous support of individual donors, local communities, and major sponsors including Cure Dale’s Duchenne, the Flatley Foundation, FibroGen, Sarepta Therapeutics, the Killian Family, Michael’s Cause, and Ryan’s Quest.
Together at Last: A Return to In-Person Summer Camp Reignites Friendships and Builds New Connections

This summer, staff, volunteers, and campers alike were thrilled to be back on the ground for in-person Camp Promise. Across five locations, the magic of camp returned and the excitement was palpable at each and every session throughout the summer.

At Camp Promise - Rockies, the Rocky Mountain scenery, crisp altitude air, and nature-filled atmosphere surrounded cabins. For some, it was like coming home. For others, a brand-new adventure.

The black cabin held most of our older campers, including Alex who is 28 years old. He says, “It is really good to be back. I enjoyed the zipline this week and hope to meet even more new people next year. My counselors were Jameel and Arturo who are really good people.”

Luke (13) did not miss a day of letting out his excitement to be back at camp. He says, “It was a lot of fun to be back and it is nice to see the new Friendship Lodge. We even got to play air hockey, pool, and video games in the game room!”

Ashley Kelliher started volunteering at camp in 2019 and has participated in many Jett Foundation events. She says, “The first time I volunteered as a counselor sparked a new level of personal growth and maturity within me. [This year] had additional precautions that both campers and staff had to take to ensure everyone’s safety, but it didn’t stop us from having an amazing week! The interpersonal relationships you build with the campers in just one week is extraordinary. Next year, I am looking forward to being reunited with old campers and introduced to new ones!”

In her first Camp Promise experience, Jett Foundation’s Director of Community Engagement Alexia Tinsley wore the hats of camp nurse and Chief Prank Approver. Alexia says, “Being back at summer camp was such a heartwarming experience. My first time at Camp Promise was like nothing I’ve experienced before. The energy, mentorship, and sense of community I experienced was life-changing and gave me a new perspective of the camp experience, as well as both teenagers and young adults living with neuromuscular diseases.”

After a long-awaited opportunity, Jett Foundation’s Executive Director Eric Snyder attended his first in-person camp since being appointed in 2019. He says, “Camp Promise Rockies provided me with a very important experience, as well as both teenagers and young adults living with neuromuscular diseases.”

In addition to our three traditional week-long camp programs they attended as children. “Camp Promise Retreat brought to life the goal of supporting young adults both at camp and in their everyday life,” says Jackie, Director of Camp Promise Retreat – Ohio. “We had opportunities to learn more about disability services and support, socialization, coping skills, mental health, and hobbies – all mixed with a lot of camp fun. The connections made are the foundation for the future.”

Throughout the retreats, campers made cabin mailboxes, decorated cabin flags, created candles, tie-dyed shirts, and painted pictures. To add to our colorful experience, campers at the Arizona retreat dressed in all white clothing and dumped colored powder onto each other during a color war.

Arizona camper Whitney shares, “My favorite thing about camp was being back with all my friends that I’ve made over the years and being able to catch up and see how they’ve been.”

"There are never enough words to explain how much Camp Promise means to me. It’s my home away from home, my second family, my safe place.” - Molly, camper

The development of Camp Promise Retreat emerged from an unmet need observed by Jodi Wolff and Jackie Simic-Recker in their local camp communities in Arizona and Ohio. Both camp directors witnessed the challenges experienced by older campers and parents as they aged out of the camp programs they attended as children.

"Camp Promise Retreat brought to life the goal of supporting young adults both at camp and in their everyday life," says Jackie, Director of Camp Promise Retreat - Ohio. “We had opportunities to learn more about disability services and support, socialization, coping skills, mental health, and hobbies – all mixed with a lot of camp fun. The connections made are the foundation for the future.”

84 volunteers across our five in-person locations and virtual camp programs

Camp Promise is made possible by our generous donors:
Ambassadors at Work: Finding Community and Fostering Supportive Friendships

As the extra eyes, ears, and voices of Jett Foundation, our Community Ambassadors serve a critical role in assisting our program team to support families nationwide. Ambassadors are crucial to helping us disseminate important information about programs. They help connect families in their circles to the right resources and encourage others to get involved to reap the full benefits of Jett Foundation’s supportive network. This year, our 31 ambassadors from 20 different states worked hard to spread awareness and engage families in their communities and beyond.

These are just some of the ways Community Ambassadors propelled our mission.

Stephanie Howie  
Mom & Advocate
- Completed the Virtual Falmouth Road Race on Team Jett
- Shared her diagnosis story and journey with pharmaceutical employees and at local speaking engagements
- Spoke in local Rare Disease Day event
- Regular participant in our monthly parent support group

Diana Johnson  
Mom & Advocate
- Joined Jett Foundation’s Board of Directors
- Panelist for Jett’s Women in Duchenne panel on World Duchenne Awareness Day
- Participated in Jett’s National Challenge

Anthony DeVergillo  
Individual Impacted by Duchenne & Advocate
- Panelist for Jett’s Thriving with Duchenne Rare Disease Day panel
- Worked with Camp Promise staff to implement new virtual camp digital environment technology
- Wrote a guest series—Rare Day in My Life—on Jett’s blog

Perlita Hains  
Mom & Advocate
- Co-facilitator of Jett Foundation’s Porch Nite Programming
- Enthusiastic advocate of Jett programs and mission
- Participates in speaking engagement sharing her journey
- JettRide host and coordinator
- Mentor to Accessible Vehicle Fund participants

Jessica Campbell  
Mom & Advocate
- Co-facilitator of Jett Foundation’s Porch Nite Programming
- Mentor to Accessible Vehicle Fund participants
- Participated in Jett Foundation’s National Challenge
- Engaged community online through social media

“This thank you for doing these. Moms like us... sometimes it’s hard to relate to our circle of friends and family no matter how much they love us and try to support... unless it’s your child, you can’t know.” - Virtual Porch Nite Participant
Empowering through Education: Workshops and Webinars Give Families Tools to be Their Own Best Advocates

Our Community Webinar Series featured thirteen online events exploring a variety of topics to help better prepare for and tackle Duchenne.

The series welcomed guest speakers from different disciplines and areas of expertise throughout the year’s 19 webinars. Speakers covered a variety of educational topics including:

- Anxiety, Behaviors, and Stress Management,
- Care Considerations When Moving States,
- Non-surgical and Surgical Management of the Tight Foot and Ankle,
- Higher Education,
- Respiratory Management,
- Vocational Rehabilitation & Driving with Duchenne,
- Navigating the Accessible Vehicle Fund,
- and more.

In addition to webinars, we were thrilled to be back in person for three traditional Family Workshops in California, Colorado, and Texas. Workshops at each location brought together families, industry partners, clinicians, and advocates for a day of educational programming, resources, and connections.

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Who requested Welcome Packets?

- Family 75%
- Medical Professional 8%
- Individual Impacted by Duchenne 6%
- Pharmaceutical Representative 5.5%
- Friend 3.5%
- Teacher 1.5%
- Advocacy Organizations >1%

Who requested Welcome Packets?

256 registrants attended our webinars live

500+ viewers watched webinar recordings

152 total registrants had never attended a previous Jett webinar

30% of Family Workshop registrants were attending to learn more about clinical trials

“Starting the Conversation: Educational Welcome Packets

Our Welcome Packets provide crucial educational materials to newly diagnosed families, caregivers at every stage of their journey, teachers, and clinicians that are seeking more guidance on Duchenne, support, and resources. Thanks to the generosity of our program sponsors, packages are provided free to community members upon request and filled with the following:

- Jett Foundation brochure and welcome letter,
- Support letters from the community,
- Health insurance, clinical trial, and classroom guides,
- Standards of care and medical durable equipment booklet,
- Dan and DMD children’s book,
- PJ’s Protocol Cards,
- and more.

This year, Jett Foundation mailed over 200 packets to individuals in 33 states, as well as a handful of international requesters.

“Thank you for the support and opportunities!”

“Thank you for the resources. I look forward to learning more about Duchenne and ways to support children with this diagnosis at school.”

“My nephew was recently diagnosed with DMD and I’m just doing as much research as I can to help me better understand it and so I can be as much help to him and his mother (my younger sister) as I can. Thank you for providing an amazing service!”

“I requested the Welcome Packet so I can show DMD families what resources you provide initially, and encourage them to engage further with your organization. Thanks!”

“I have a student this year with DMD and I want him to feel welcome and for his peers to understand what he is going through and to be kind.”

“I am a Physical Therapist Student and my supervising Clinical Instructor and I are super interested in learning everything we can to give our patient the best quality of life.”

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“"The webinar topic was very interesting and the speaker did a great job explaining things in an easy-to-understand format.”

“I liked that the family perspective presentations were an official part of the Family Workshop agenda - I think it shows that their stories are valued and just as important as the clinical presentations.”

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Thriving with Duchenne: Rare Disease Panel Showcases Individual Achievements

On February 28, 2022, Jett Foundation hosted a virtual event to celebrate Rare Disease Day 2022. This was Jett Foundation’s 5th annual Rare Disease Day, which welcomed speakers from the Duchenne and rare disease communities in a panel discussion themed “Thriving with Duchenne.”

Moderating the event was Kyle Bryant and Sean Baumstark, the hosts of Two Disabled Dudes podcast. Kyle and Sean are both adults living with a rare disease called Friedreich’s ataxia (FA). Jett Foundation is very grateful for their presence at Rare Disease Day.

Also speaking at the event was Elijah Stacey, Anthony DeVergillo, Leslie Crowley, and Guðjón R. Óskarsson, who are all men living with Duchenne muscular dystrophy.

Anthony, 29, feels he has come a long way since first being diagnosed with Duchenne: “From a young age, I always had trouble being social, fitting into social groups, or trying to hang out with my friends. Now, I am an optimistic and outgoing person and I have found other ways to be social and get involved through the virtual world and technology.”

Over 200 people from across the nation joined the panel discussion and even more participated on social media, sharing our infographics leading up to Rare Disease Day.

The team at Jett Foundation would like to extend our gratitude to all of our sponsors for supporting this event and making our Rare Disease Day 2022 celebration possible. Please join us in thanking Antisense Therapeutics, Avidity Biosciences, Casimir, Dyne Therapeutics, Edgewise Therapeutics, Entrada Therapeutics, FibroGen, Hallymaco, N3 Pharma, PepGen, Pfizer, RegenXBio, Sarepta Therapeutics, Solid Biosciences, Stealth Biotherapeutics, and Ultragenyx Pharmaceutical.

“As the head of patient advocacy for a biotech company, I will typically arrange for a guest speaker and group activity for our team to recognize Rare Disease Day. But when I saw the lineup of speakers you’d secured, I abandoned my plan for an independent event and knew our time would best be used tuning into your panel discussion.” - Event Participant

Broadening Reach: Jett Celebrates World Duchenne Awareness Day

On September 7, 2022, Jett Foundation hosted its seventh annual World Duchenne Awareness Day (WDAD) celebration to recognize those in the Duchenne community. To bring awareness, fundraising, and joy into the homes and offices of our community, Jett Foundation offered WDAD Awareness Boxes that included items for participating online, raising awareness for Duchenne, and showing Jett Foundation spirit.

“Jett Foundation’s World Duchenne Awareness Day celebration is very important for being able to broaden our reach nationally and globally,” says Eric Snyder, Jett Foundation’s Executive Director.

Our first interactive event of the day was a Women in Duchenne panel discussion, which featured women carriers, women caregivers, and women in the clinical care setting sharing their experiences. Speakers included: Dr. Migwé Monday, Director of the Neuromuscular Program at Nicklaus Children’s Hospital; Trina Stelly, a Duchenne carrier, mother to a family impacted by Duchenne, and Jett Foundation Ambassador; and Diana Johnson, a mother to three children, one with Duchenne, and Jett Foundation Ambassador.

The panel was facilitated by Mindy Cameron, a Duchenne carrier, long-time Patient Advocate, and the mother of an individual who lives with Duchenne. Mindy says, “Every day, we hear of Duchenne being a disease that affects males. I think we are all trying to move away from that strict definition. The fact that there are now dedicated clinics for women like us has been really significant.”

On the topic of raising awareness about both male and female Duchenne carriers, Trina Stelly shared, “When I started doing more research, I found that there were other women showing my same signs and symptoms. I try to participate in as many research studies as possible so that we can find out the answers to our questions. As hard as it is, [females] need to take the time to look into themselves.”

In the afternoon, the Jett Foundation team get up out of their chairs to dance with the community in our Dancing for Duchenne event.

Facilitated by Hulda Björk and her son, Ægir Þór, this dancing session was one of many global stops that the duo has made to raise awareness for Duchenne muscular dystrophy.

Hulda says, “Having a son with Duchenne, I wanted to do something good. I needed to have a purpose and meaning to all of this, so I started advocating and doing little things on social media. One day, we posted a dancing video. It became a big thing, and now we are Dancing for Duchenne!”

The community’s energy stayed high through the evening as we concluded our day with an interactive trivia game. Containing two teams composed of individuals who all come from different areas in the Duchenne space, this game brought a fun and competitive battle of answering miscellaneous trivia questions about Duchenne and pop culture.

Our 2022 WDAD celebration would not be possible without the many individuals, families, partners, and teams around our community. We can’t thank you enough for your participation!
Solidifying Our Future

Fox Rock Foundation Gifts One Million Dollar Endowment to Jett Foundation

Jett Foundation is thrilled to share that we have received a $1 million endowment gift from Rob and Karen Hale through the Fox Rock Foundation, the couple’s family foundation established alongside their adult children, Trevor, Thomas, and Brett. The funds will be used at the foundation’s discretion to sustain and expand the organization’s direct-service programming and advocacy for individuals and families impacted by Duchenne muscular dystrophy.

“This is a wonderful moment for Jett Foundation and we are honored by this incredible act of philanthropy from the Fox Rock Foundation and the Hale family,” said Eric Snyder, Executive Director.

Together with the support of donors like the Hale family, in this past year alone, Jett Foundation provided over 200 individuals with week-long in-person and virtual summer camp opportunities, educated over 500 community participants through its educational webinar series, provided support and grief management to over 190 moms, grandmothers and caregivers through its monthly parent support group program, and provided accessible vans and emergency grants to dozens of families in need through its Jett Giving Fund.

“Over the past few years we have had the privilege of supporting Jett Foundation and coming to know the deep impact of Jett Foundation’s family-focused programs,” says Karen Hale. “We are excited to help ensure the future of Jett Foundation as they aim to serve as many families in the Duchenne community nationwide as they can.”

The gift comes at a pivotal moment in Jett Foundation’s history, with the organization having recently undergone an extensive three-year strategic plan.

“Our new strategic plan maps out a path forward to advance the impact of the programming our community loves and depends upon while focusing on reaching more people impacted by Duchenne nationwide,” says Snyder. “The Hale’s gift will help make this vision a reality, strengthening today’s programs while solidifying our future.”

Thanks to the Hales’ philanthropy, and the ongoing generosity of Jett Foundation friends and supporters, the future is bright for our long-term ability to provide transformational programming to the Duchenne muscular dystrophy community.

Team Jett Takes Falmouth

Jett Foundation Athletes Race and Raise Awareness Along the East Coast and Beyond

Team Jett is Jett Foundation’s endurance athletic team that participates in many events each year to raise funds and awareness for Duchenne muscular dystrophy. In 2022, our team participated in the Cohasset Triathlon, Falmouth Road Race, Hopkinton Title 9 Triathlon, Rodman for Kids bike ride, and a National Challenge hosted by Jett Foundation.

Our Jett red colors shined bright through the historic streets of Falmouth, MA at the 50th running of the Falmouth Road Race. Thirteen Team Jett athletes completed the 7-mile course on race day, along with two virtual runners who traversed their own course in Plymouth, MA. With streets filled with spectators and supporters, it was impossible to not have a successful day.

Jett Foundation’s own Marketing and Communications Manager Zac Bentley joined our 15-person lineup to raise funds and awareness for Duchenne. With the support of family and friends, Steph and James Howie and James Bustamante hosted their own road race in Plymouth, Massachusetts while raising an abundance of funds and awareness for Duchenne. With the support of family and friends, Steph and James completed a 7-mile loop across the Plymouth waterfront on August 14, 2022 in honor of their two-year-old son Gio, who has Duchenne.

“Our son, Giovanni, was diagnosed at just 4 months old with Duchenne muscular dystrophy,” says Steph. “While this news was absolutely life shattering, we were introduced to Jett Foundation and they have been such an incredible support system for our family. They have provided us with so many tools we need to fight this horrific disease and a community to help us feel like we are not on this journey alone.”

She also says, “We were thrilled to have the chance to be part of the Falmouth Road Race Team Jett runners and help raise funds to continue helping families just like ours. It was such a great experience walking the 7 miles through Plymouth... An experience we will never forget and look forward to doing again.”

Kyla Collette and Alex Tuckerman also finished strong in Falmouth where they raced for their son, Walker, who was diagnosed with Duchenne just after his first birthday.

Special thank you to our 2022 Team Jett sponsors: Shea Concrete Products, Solid Biosciences, Yale Appliance, BayCoast Bank, Quincy Mutual Group, and South Shore Bank.

Members of Team Jett raised nearly $23,000 through the Falmouth Road Race in support of Jett Foundation’s programs and services that assist individuals and families impacted by Duchenne muscular dystrophy.

Steph Howie and James Bustamante hosted their own road race in Plymouth, Massachusetts while raising an abundance of funds and awareness for Duchenne. With the support of family and friends, Steph and James completed a 7-mile loop across the Plymouth waterfront on August 14, 2022 in honor of their two-year-old son Gio, who has Duchenne.

“Over the past few years we have had the privilege of supporting Jett Foundation and coming to know the deep impact of Jett Foundation’s family-focused programs,” says Karen Hale. “We are excited to help ensure the future of Jett Foundation as they aim to serve as many families in the Duchenne community nationwide as they can.”

The gift comes at a pivotal moment in Jett Foundation’s history, with the organization having recently undergone an extensive three-year strategic plan.

“Our new strategic plan maps out a path forward to advance the impact of the programming our community loves and depends upon while focusing on reaching more people impacted by Duchenne nationwide,” says Snyder. “The Hale’s gift will help make this vision a reality, strengthening today’s programs while solidifying our future.”

Thanks to the Hales’ philanthropy, and the ongoing generosity of Jett Foundation friends and supporters, the future is bright for our long-term ability to provide transformational programming to the Duchenne muscular dystrophy community.
Thank you to our donors

Our donors are what makes it possible for Jett Foundation to empower families and fight Duchenne. Thank you to the people and organizations who provided a contribution of $1,000 or greater during FY22 through financial gifts, in-kind donations, or peer-to-peer fundraising.

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Amanda McAride
Amy Brantner
Angela Straenner
Anne Fried
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Avidity Biosciences
Bay State Federal Savings Charitable Foundation
BayCoast Bank
Bernard Reynolds
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Tyler Lafferty
Ultrapenny Pharmaceutical Inc.
Vertex Pharmaceuticals
Walking To End Duchenne
Walt and Nita Simmerman
Winifred Belmont

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FY22 Financials

<table>
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<tr>
<th></th>
<th>Revenue</th>
<th>Expenses</th>
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<tr>
<td>Corporation &amp; Foundation</td>
<td>$1,604,750</td>
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<tr>
<td>Individual Donations</td>
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<td>Programs &amp; Events</td>
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<td><strong>Total Support</strong></td>
<td><strong>$2,305,217</strong></td>
<td><strong>$2,020,527</strong></td>
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</table>

Use of Funds

- **Program Services**: 78%
- **Fundraising**: 16%
- **General & Administrative**: 12%

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Thanks to everyone who supported our mission this year!
STAFF

**Eric Snyder**, Executive Director

**FINANCE & HR**
Allison Caputo
Kelly MacGrath

**PROGRAMS**
Niki Benfield
Meghan Houston
Julie LeClair
Alexa Tinsley

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**Executive Board**

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Casimir

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Alfred Jackson, Vice-Chair
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Robert Stubbs, Careport Health

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Susanna Doyle
Caitlin Hughes

**MARKETING & COMMUNICATIONS**
Zac Bentley
Gabriella Costa
Cori Dubois