The Parkinson’s Foundation makes life better for people with Parkinson’s disease by improving care and advancing research toward a cure. In everything we do, we build on the energy, experience and passion of our global Parkinson’s community.

$24M invested in Parkinson’s research and clinical care in 2023.

On September 9, 2023, our Southwest Chapter hosted PD 101 in Tucson, AZ. More than 100 attendees heard from Cynthia Reed, MD, of the Parkinson Wellness Clinic and enjoyed a movement break with PWR!

Board member Christina Weaver Jackson (left) alongside Halston Mathis (second from right) at the Move & Mingle event in Washington, DC, on March 16, 2023.

On January 30, 2023, 73 people attended our Promotores de Salud day-long training workshop on Parkinson’s, research and resources in Chicago, IL.

On September 9, 2023, our Southwest Chapter hosted PD 101 in Tucson, AZ. More than 100 attendees heard from Cynthia Reed, MD, of the Parkinson Wellness Clinic and enjoyed a movement break with PWR!
In 2023, we dedicated $24 million to Parkinson’s research and clinical care. We are seeing unprecedented momentum in our global genetics study, PD GENEration: Mapping the Future of Parkinson’s Disease. We have already provided genetic testing and counseling to 13,600 participants, at no cost, and this pace will continue in the year ahead as we expand to new areas.

We published an eye-opening study this year about critical gaps and disparities in access to Parkinson’s disease (PD) care in the U.S. One of the key findings is that only 9% of people with Parkinson’s received care from movement disorders specialists, expertly trained neurologists who can recognize PD’s distinct nuances and tailor treatments to each patient. These findings underscore the need for greater outreach to general neurologists and other providers to better understand their needs in caring for people with Parkinson’s.

Two years ago, we launched our Reach Further fundraising campaign to invest an additional $30 million to accelerate progress on Parkinson’s research, improve care and increase access to quality-of-life programs. We are pleased to share that we have raised more than $16 million and are on track to meet our goal well before 2025 thanks to our many generous donors, including the Krupp Smith Family.

“We were lucky our dad had access to great care, support and educational resources. By supporting the Parkinson’s Foundation, we are expanding access to care, supporting research and honoring our parents’ legacy,” said Bobby Smith and Gary Smith, Jr. who honored their late father, Gary Smith Sr., with a gift to the Reach Further campaign.

I think you will see for yourself in this report that we are accelerating our efforts in research, care and education. Thank you to the clinicians who care for people with Parkinson’s disease every day, to the scientists who provide hope for breakthroughs and a better future, to our volunteers for their energy and tireless efforts, and to our donors for their faithful support of our mission.

Sincerely,

John L. Lehr
President and Chief Executive Officer

The cast of the Broadway play A Beautiful Noise presented John Lehr with a check for $500,000 on April 11, 2023, to help make life better for people with PD.

Last year our People with Parkinson’s Advisory Council met in Miami, FL, to help guide Foundation priorities and programs.

Our Carolinas Chapter premier golfing event, Playing for Parkinson’s, at Spring Island, SC, raised $325,000 and brought together local individuals and businesses.
$449 million invested in Parkinson’s research and clinical care since 1957.

678 scientists received Parkinson’s Foundation funding since 2010.

13,600 people with Parkinson’s enrolled in PD GENERation, to receive genetic testing and counseling at no cost.
Although an estimated 10 million people live with Parkinson’s globally, research remains severely underfunded. At the Parkinson’s Foundation, we are committed to closing the funding gaps in research by providing essential grants, furthering drug discovery, and advancing our global genetics study.

A recent Parkinson’s Foundation study found that nearly 90,000 people are diagnosed with Parkinson’s every year in the U.S. — this number is 50% higher than previously believed. To understand Parkinson’s and how to better help people, we need to know how many people live with this disease, which is why studies like this one are imperative.

In 2023, we awarded $3.5 million in research grants to 32 scientists. Through grants and fellowships, we fund scientists who are devoted to deciphering this disease and finding new ways to slow its progression. We back high-risk research that typically does not receive federal funding, so that we can advance the research that leads to high rewards.

We invested $3 million toward the Parkinson’s Virtual Biotech, in partnership with the Parkinson’s UK. This initiative is funding the development of 13 potential drugs that either address symptoms or aim to slow, stop, or prevent the disease altogether, getting us closer to delivering life-changing treatments in years, not decades.

Right now, pharmaceutical companies are developing drugs that slow PD through targeting specific genetic mutations — some are already recruiting for clinical trials. However, it can take years to fill these trials. Not only does PD GENEration: Mapping the Future of Parkinson’s Disease help accelerate clinical trials, and their results, but this data is immediately shared with researchers to develop a greater understanding of the genetic link to Parkinson’s.

In 2023, PD GENEration:
• Provided genetic testing and counseling to 13,600 people with Parkinson’s, at no cost.
• Found that up to 12.7% of people with Parkinson’s have a genetic link to the disease.
• Dedicated efforts to recruiting diverse applicants and reaching priority populations.

Our research fellowships and early career awards support and train young investigators. The Parkinson’s Foundation Launch Award enables postdoctoral researchers to complete needed mentored training and transition promptly to independent research careers in Parkinson’s.

“The Parkinson’s Foundation Launch Award will have a profound impact on my career and abilities to continue Parkinson’s research. Opening a new laboratory as an early career scientist will be an exciting adventure. I will face many challenges, but this grant will greatly facilitate my success. Thanks to all Parkinson’s Foundation supporters for helping us move the research forward.”

While Parkinson’s is complex, our research strategy is simple — accelerate Parkinson’s research on every level.

Learn more at Parkinson.org/Research.
11,063 emails sent to people in the PD community with resources and referrals from our Helpline in 2023.

8 new specialized PD centers joined our Global Care Network in 2023, with 7 in the U.S. and our first-ever in Japan.

9,000+ credits given across 12 health professional disciplines through our online PD courses in the Learning Lab.
Regularly seeing a neurologist can improve the lives of thousands of people with Parkinson’s disease each year. Our recent care access study, based on Medicare data, found that 50% of people with Parkinson’s see a general neurologist. This study serves as our blueprint for improving access to PD care. We are focused on reaching people where they are most likely to receive care and the professionals who provide it. We are amping disease-specific training, resources and collaborating with community partners. This brings us closer to improving access to care across all demographic and population groups.

To bring Parkinson’s care to more people, we designate medical centers that know how to treat this complicated disease. These centers utilize a specialized team-based approach and provide community outreach. In 2023, we designated eight new centers as part of our Global Care Network, with the first-ever designations in Wisconsin and Washington, and the country of Japan.

In 2023 we expanded our network to include Community Partners in Parkinson’s Care, a membership program for senior living communities and home care agencies. Through this program, more than 70 sites have staff trained in providing Parkinson’s care.

Even with an aging population and Parkinson’s on the rise, most healthcare professionals do not receive in-depth Parkinson’s care training. With support from the CVS Health Foundation, we launched a new Education Series for Community Providers — an accredited free series of online courses designed to improve treatment and outcomes for people with Parkinson’s.

From designating care centers to offering courses for healthcare professionals and the Parkinson’s community, we believe in helping everyone understand the best-practices in Parkinson’s care. All of our tools are designed to empower people with Parkinson’s to advocate for the care that will lead to a better quality of life.

When people with Parkinson’s visit one of our Global Care Network centers, they can expect high-quality care from professionals who know the latest in treatments and research. Expanding this network is just one way we are increasing access to care.

The Stanford Movement Disorders Center is deeply honored to be designated as a Parkinson’s Foundation Center of Excellence. Through patient care, education and research, our providers and staff embrace the Foundation’s mission to make life better for people with Parkinson’s.

Explore how we are accelerating access to Parkinson’s care at Parkinson.org/FindingCare.
Nearly $1 million awarded in community grants benefiting Parkinson’s programs across 35 states.

123,085 people registered for in-person and virtual PD events — reaching 42 countries and 50 U.S. states.

81,060 educational books and 386,200 fact sheets distributed.
Learning to live well with Parkinson’s disease is an ongoing endeavor, which is why our resources and events are designed to empower those living with this disease, care partners and healthcare professionals.

Ensuring diversity and inclusivity in Parkinson’s research is critical to accelerating breakthroughs toward a cure. In 2023, we partnered with Morehouse School of Medicine to expand genetic research in the Black community, making Morehouse the first historically Black institution of medicine to become a PD GENEration research site. Through PD GENEration, we make no-cost genetic testing and counseling accessible to everyone with PD. This includes local education opportunities, including our in-person research advocate training in Atlanta, GA, attended by 33 advocates from the Black community.

With 63.6 million people in the U.S. of Hispanic and Latino origin (according to the U.S. Census Bureau), we are dedicated to reaching the Spanish-speaking Parkinson’s community. We launched more than 50 new Spanish pages on Parkinson.org and created the Parkinson’s Foundation Spanish Advisory Group to help us identify educational gaps and design educational programs for healthcare professionals. To provide peer-to-peer education and support for the Hispanic community, we hosted four Promotores de Salud training workshops with 127 attendees. Our PD GENEration genetics study also extends to the Spanish-speaking community, providing genetic testing and counseling to 1,060+ Latino participants, in English and Spanish.

Access to information is vital for people living with Parkinson’s, and those who love them. We funded 71 community grants across 35 states — from local Parkinson’s non-contact boxing classes to support groups. We also hosted 196 national and community-based education programs with 123,085 registrations. Every one of our programs and events are designed to fill an unmet need for the Parkinson’s community.

There is an unprecedented need for community-related programs that benefit people with Parkinson’s and their care partners. From exercise classes to wellness events, our community grants aim to further the health, wellness and education of people with Parkinson’s across the U.S.

“The support we have received from the Parkinson’s Foundation for our What’s Up, Doc? program has enabled us to have high touch opportunities with people in the first year following a diagnosis. This program brings together the newly diagnosed in the local community to get them involved. Early intervention, education and inclusion in our community will enable those who are newly diagnosed to live a better life with the disease.”

Accelerate your Parkinson’s community and find empowering resources and events at Parkinson.org/Resources.
2023 in Numbers

302,582 people received vital counseling from specialists on our toll-free Helpline, 1-800-4PD-INFO since 1998.

204,440 people with Parkinson's treated in our Global Care Network — the largest global network of medical centers that specialize in Parkinson's care.

3,150 volunteers helped us make life better for people with PD across all 17 of our chapters.

$449 million invested in Parkinson's research and clinical care since 1957.

678 scientists received Parkinson's Foundation funding since 2010.

787 community-based programs have received $10.4 million in funding since 2011.
50 new webpages with critical information about Parkinson's on Parkinson.org/Español.

13,600 people with Parkinson's received a genetic test and counseling session through our Parkinson's study, PD GENEration.

6 million people visited Parkinson.org, the go-to online Parkinson's resource, in 2023.

117,000 podcast episodes downloaded in English and Spanish, in 2023.

12.7% of participants have a genetic link to PD (much higher than the originally believed 5-10%).

74% of participants have never participated in a PD research study prior to PD GENEration, opening doors for people to join other research studies and advance the field.
ACCELERATING HOPE

Halfway into our Reach Further campaign, we have received 6,611 donations that help us advance Parkinson’s research, improve care and increase access to quality-of-life programs. Funds raised in 2023 allowed us to expand our genetics study, PD GENERation, invest in drug discovery and development through the Parkinson’s Virtual Biotech, fund community programs and expand our Global Care Network. Thank you to our generous supporters!

Help us reach further at Parkinson.org/Reach.
NATIONAL CORPORATE SPONSORS

The Parkinson’s Foundation gratefully acknowledges our national corporate sponsors who actively support mission-critical programs and research initiatives.

Amneal Specialty Pharmaceuticals

Kyowa Kirin

Supernus

ACADIA Pharmaceuticals, Inc.

CVS Health Foundation

Biogen, Inc.

Boston Scientific Corporation

AbbVie

Medtronic

Abbott Laboratories

Merz Therapeutics

Avion Pharmaceuticals, LLC

ACORDA Therapeutics

Sunovion Pharmaceuticals, Inc.

Neurocrine Biosciences, Inc.

Genentech Inc., Member of the Roche Group

Preval Therapeutics

Sage Therapeutics

Right At Home, LLC

Neal and Sherrie Savage

Family Foundation

The Scarlett Family Foundation

Estate of Jane M. Schaal

Carl S. Schneider

Fred and Nancy Schumacher

Dr. Vilasini Shanbhag*

Eddie and Jo Allison Smith

Family Foundation

Estate of Kathleen A. Steinwehr

William Harris and Andrea Sussman*

Daniel A. Swiglo

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Jeff Fisher*

James R. Welty

Anne Wojcicki Foundation

Yonkers Contracting

Company, Inc.

Sherwin and Sheri Zuckerman

*denotes multi-year pledge donor
Donor Spotlight: How Two Families Help Accelerate Parkinson’s Research

Gurley Family Amplifies Impact Through Funding Innovative Research Studies

“When I learned about the type of research the Parkinson’s Foundation is funding, studies that might not get funded through a traditional process, that really spoke to me,” said Bill Gurley.

For Bill and Amy Gurley, finding a cure to Parkinson’s disease (PD) is personal. After watching Bill’s mother, Lucia, battle PD, they felt compelled to help solve Parkinson’s through funding critical research. “There’s so many aspects of Parkinson’s you can’t even begin to understand,” Amy said. “It creates a crisis within a family. Lucia had such a kind heart and to watch her degenerate was the saddest part. She is what drove us to figure out how to be a servant of this.”

Bill and Amy generously funded the 2023 Impact Awards, supporting six innovative research studies, each with the potential for a breakthrough. “I’ve been a venture capitalist my whole life and there’s a saying that risk and reward are correlated,” said Bill. “If you want to increase the optionality of outsized impact, or discovery, you have to be willing to invest in things that might not be obvious.

One of the things that energize Bill and Amy most about Impact Awards is that each recipient is aiming to find a root cause of Parkinson’s, which could result in a monumental impact, including a cure for PD.

“‘When Parkinson’s touches your family, you see how powerful and difficult it is to go through,’” Bill said. “I can’t imagine the impact if you don’t have access to resources. I hope that through our efforts, and through the support of other people investing in research, we can find a solution that has more of a permanent impact on this disease.”

Gary and Bobby Smith Honor Their Parents’ Commitment to Parkinson’s Research

“He was patient number one at University of Kansas (KU) Medical Center,” said his son Bobby Smith. “When his doctor shared the initiative with him, he was on board immediately. He was always ready to help whenever he was able — he wanted something different for other people facing this disease.”

Now, his sons, Gary Smith, Jr., chairman of the Krupp Smith Family Foundation, and Bobby Smith, president, are proud to honor their late father by supporting the Reach Further campaign and expansion of PD GENEration. Thanks to their generosity, the Parkinson’s Foundation Krupp Smith Family Foundation Center of Excellence and Gary A. Smith PD GENEration Site at KU Medical Center were named in their family and father’s honor.

“When a family member is diagnosed with Parkinson’s, it is hard to connect to the right resources and find the best care,” Gary Jr. said. “If we can help improve understanding of the disease and access to care for other people, that’s a big deal and it is really important to us.”

The passion their father had for research continues to inspire Gary Jr. and Bobby. “We were lucky our dad had access to great care, support and educational resources,” Bobby said. “My parents were so generous and always looking for ways to help others. By supporting Reach Further, we are expanding access to care, supporting research and honoring our parents’ legacy.”

There are many ways you can support the fight against Parkinson’s. Visit Parkinson.org/HowToHelp.
Parkinson’s Foundation
Legacy Society
The Parkinson’s Foundation recognizes and honors all Legacy Society members who support the Foundation in their wills, trusts, life income gifts, retirement plans and other planned gifts. Legacy Society members leave meaningful, lasting legacies for future generations.

Lester Nathan Leaves a Legacy Through His Estate Plan

“"The number of people with Parkinson’s is increasing, and the Foundation uses a lot of its funds for essential research to determine its causes and find a cure. I believe the Parkinson’s Foundation is a well-run organization and that anyone, directly or indirectly, affected by Parkinson’s should consider including the Foundation in their estate plans because it is a good way to honor their loved ones’ memories.”

– Lester Nathan

Before his aunt, Mechtild Eisner, was diagnosed with Parkinson’s, Lester Nathan was not aware of the Parkinson’s Foundation. Since learning more about PD and the resources the Foundation offers people living with the disease, Lester was inspired to add the Parkinson’s Foundation to his estate plans, making him a member of our Legacy Society. “I want to make sure that part of my estate will continue to help the charities I support, including the Parkinson’s Foundation,” Lester said. “In a way, this bequest also helps me continue my legacy and honor my aunt.”

After learning more about Parkinson’s and other neurological conditions, Lester was inspired by the Foundation’s efforts to support research devoted to understanding what causes PD, which will lead to new treatments.

As a Legacy Society member, Lester supports advances in research and resources, which help people live better with Parkinson’s disease every day.

Learn more about the Legacy Society and planned giving options at Parkinson.org/PlannedGiving.

In Memoriam

Lucien J. Côté, MD

We remember the legacy of Lucien J. Côté, MD, one of the most respected and beloved Parkinson’s doctors in the U.S. For more than 60 years, Dr. Côté served on the neurology faculty at Columbia University Irving Medical Center, a Parkinson’s Foundation Center of Excellence. In 2014, he received our Lifetime Achievement Award.

Richard “Dick” D. Field

We honor the loss of emeritus board member Richard “Dick” D. Field, who served as a board member from 2013 to 2021. Co-founder of LendingTree and founder of Richard D. Field Associates, Dick was a financial services industry leader and mentor. After his Parkinson’s diagnosis, he served the Foundation as a passionate advocate and fundraiser, playing an instrumental role in funding ground-breaking initiatives in PD care and research.
Moving Day, A Walk for Parkinson’s

At Moving Day, we honor loved ones, raise funds, fight Parkinson’s disease and celebrate movement — proven to help manage Parkinson’s symptoms. Join a team, become a sponsor or volunteer to help beat Parkinson’s at one of our 53 events taking place in 2024.

After being diagnosed with Parkinson’s, Kathleen Ortiz attended Moving Day looking for hope. Today, she volunteers to provide hope to others. “In order to give the next person who will be diagnosed more hope than I had, we have to get out there and walk, raise money and talk about Parkinson’s. Volunteering with the Parkinson’s Foundation has taken me into a direction I never thought I would go. At Moving Day, I feel accepted — and that feels so good. I will do anything I can to help someone else find that comforting feeling.”

Move with your community at Moving Day and help make life better for people with Parkinson’s. Find your nearest Moving Day at MovingDayWalk.org.

$4.8M Raised in 2023
46 Events in 2023
$44.8M Raised since 2011
13,000 Participants in 2023

Top 10 Moving Day Participants
Amount Raised in 2023

$49,428 Sherwin Zuckerman Chicago
$39,950 Andrew Albert Chicago
$33,047 Christine Howard Twin Cities
$25,043 Susan Brown Atlanta
$19,420 John Kolaya New Jersey

$19,077 Jan Wuliger Tampa Bay
$18,305 Laura Morris Chicago
$15,435 Carey Durham Dallas/Fort Worth
$14,381 Shari Wolf Chicago
$13,380 Margaret Samaha Santa Fe
Our Parkinson’s Champions are energized to help us beat Parkinson’s. Champions have raised $21 million to make life better for people with Parkinson’s. We thank all Parkinson’s Champions and recognize our top fundraisers of 2023.

### Top 10 Parkinson’s Champions

<table>
<thead>
<tr>
<th>Rank</th>
<th>Name</th>
<th>Event</th>
<th>Raised</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Karan Rai</td>
<td>World Marathon Challenge</td>
<td>$80,374</td>
</tr>
<tr>
<td>2</td>
<td>Mike Debartolo</td>
<td>Playing for Parkinson’s</td>
<td>$76,646</td>
</tr>
<tr>
<td>3</td>
<td>Roland Frankel</td>
<td>Graeme Frankel Memorial Golf Tournament</td>
<td>$70,145</td>
</tr>
<tr>
<td>4</td>
<td>Christian Spence</td>
<td>Going the Distance for PD</td>
<td>$22,200</td>
</tr>
<tr>
<td>5</td>
<td>Steven Goode</td>
<td>The Great American Scenic Byways Tour</td>
<td>$21,795</td>
</tr>
<tr>
<td>6</td>
<td>Paul Farahvar</td>
<td>Comedy Night at Zanie’s</td>
<td>$21,421</td>
</tr>
<tr>
<td>7</td>
<td>Joyce Tracy</td>
<td>Zero Mile Fun Run</td>
<td>$15,221</td>
</tr>
<tr>
<td>8</td>
<td>Carl Munyon</td>
<td>60th Birthday Golf Tournament</td>
<td>$14,992</td>
</tr>
<tr>
<td>9</td>
<td>MIG Bearden</td>
<td>Pickleball for Parkinson’s</td>
<td>$12,435</td>
</tr>
<tr>
<td>10</td>
<td>Joe Violi</td>
<td>Jimmy Violi Memorial Golf Tournament</td>
<td>$8,800</td>
</tr>
</tbody>
</table>

### Top 10 Parkinson’s Champions: Endurance Race

<table>
<thead>
<tr>
<th>Rank</th>
<th>Name</th>
<th>Event</th>
<th>Raised</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Jamie Harrison</td>
<td>2023 TCS NYC Marathon</td>
<td>$38,483</td>
</tr>
<tr>
<td>2</td>
<td>Bruce Braswell</td>
<td>2023 TCS NYC Marathon</td>
<td>$29,432</td>
</tr>
<tr>
<td>3</td>
<td>Liz Hughes</td>
<td>2023 TCS NYC City Marathon</td>
<td>$28,000</td>
</tr>
<tr>
<td>4</td>
<td>Craig Fuller</td>
<td>2023 TCS NYC City Marathon</td>
<td>$14,250</td>
</tr>
<tr>
<td>5</td>
<td>Amy Gottlieb</td>
<td>2023 Falmouth Road Race</td>
<td>$13,028</td>
</tr>
<tr>
<td>6</td>
<td>Ladd Harrison</td>
<td>2023 TCS NYC Marathon</td>
<td>$12,895</td>
</tr>
<tr>
<td>7</td>
<td>Annie Taylor</td>
<td>2023 TCS NYC Marathon</td>
<td>$11,435</td>
</tr>
<tr>
<td>8</td>
<td>Catherine Griggs</td>
<td>2023 TCS NYC Marathon</td>
<td>$10,968</td>
</tr>
<tr>
<td>9</td>
<td>Saul Qersdyn</td>
<td>2023 TCS NYC Marathon</td>
<td>$9,350</td>
</tr>
<tr>
<td>10</td>
<td>Melanie Johnson</td>
<td>2023 TCS NYC Marathon</td>
<td>$8,318</td>
</tr>
</tbody>
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### Top Parkinson’s Revolution Riders

<table>
<thead>
<tr>
<th>Name</th>
<th>Location</th>
<th>Raised</th>
</tr>
</thead>
<tbody>
<tr>
<td>Melanie Hannon</td>
<td>Chicago</td>
<td>$17,878</td>
</tr>
<tr>
<td>Deborah Lucchesi</td>
<td>San Francisco</td>
<td>$12,388</td>
</tr>
<tr>
<td>Jennifer Goldman</td>
<td>USA</td>
<td>$11,165</td>
</tr>
<tr>
<td>Chas Schaner</td>
<td>Grand Rapids</td>
<td>$11,050</td>
</tr>
<tr>
<td>Brett Pinegar</td>
<td>Salt Lake City</td>
<td>$11,010</td>
</tr>
</tbody>
</table>
As illustrated below, the Parkinson’s Foundation directed 81% of its 2022-2023 spending on mission-related activities and 19% on fundraising and management combined.*

**Total Revenue**
$48,333,484

**Total Year-End Assets**
$47,034,483

**Total Operating Expenses**
100% | $48,541,314

- **Subtotal Mission-Related**
  81% | $39,308,989

- **Fundraising**
  14% | $6,560,910

- **General & Administration**
  5% | $2,671,415

**REVENUE by Category**

- **$7,899,450**
  Special Events, Peer to Peer & All Other**

- **$9,587,354**
  Institutional Giving: Corporate & Foundation

- **$30,846,680**
  Individual Giving: Private & Bequests

**MISSION-RELATED EXPENSES by Category**

- **$11,342,739**
  Ensuring Better Outcomes through Improved Care

- **$12,573,516**
  Understanding Parkinson’s through Research

- **$15,392,734**
  Educating & Empowering People with and Affected by Parkinson’s

* Reflects Parkinson’s Foundation 990 for the fiscal year 2022-23
** Includes Investment Income
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Parkinson's Foundation Board of Directors are dedicated volunteers who guide us through their skilled leadership — invaluably setting the course towards an impact-driven mission.

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*Passed Away*