JOIN THE MOVEMENT AS WE RALLY FOR RARE

GET READY TO SHINE
Dear Nevus Outreach Community,

I am thrilled to share the inaugural edition of "The Dotted Line- A Nevus Outreach Publication." Your unwavering support and dedication have helped us forge a beautiful community, and I am proud of our progress together.

While we celebrate our achievements, it is important to acknowledge that our journey is far from complete. There is still much work ahead as we strive to reach new heights in advocacy, support, and research. With your continued commitment, I am confident that we can overcome challenges and make a lasting impact on the lives of those affected by CMN.

As the Executive Director, it is an absolute pleasure to lead this community during such a pivotal time where research interest, advocacy efforts, and community support are building to a trifecta of energy. Your enthusiasm and engagement are driving us towards a future where individuals with CMN and their families find hope, understanding, and a supportive network.

Thank you for being an integral part of our community and contributing to our mission every day. Together, we will build a brighter future for individuals with CMN and their families.

Warm regards,

[Signature]

Executive Director
Nevus Outreach

A NOTE FROM LAUREN

"While we celebrate our achievements, it is important to acknowledge that our journey is far from complete. There is still much work ahead as we strive to reach new heights in advocacy, support, and research."

Dear Nevus Outreach Community,

I am thrilled to share the inaugural edition of "The Dotted Line- A Nevus Outreach Publication." Your unwavering support and dedication have helped us forge a beautiful community, and I am proud of our progress together.

While we celebrate our achievements, it is important to acknowledge that our journey is far from complete. There is still much work ahead as we strive to reach new heights in advocacy, support, and research. With your continued commitment, I am confident that we can overcome challenges and make a lasting impact on the lives of those affected by CMN.

As the Executive Director, it is an absolute pleasure to lead this community during such a pivotal time where research interest, advocacy efforts, and community support are building to a trifecta of energy. Your enthusiasm and engagement are driving us towards a future where individuals with CMN and their families find hope, understanding, and a supportive network.

Thank you for being an integral part of our community and contributing to our mission every day. Together, we will build a brighter future for individuals with CMN and their families.

Warm regards,

[Signature]

Executive Director
Nevus Outreach
As of the close of the fiscal year 2023, Nevus Outreach reports a total gross revenue of $210,819, marking a significant achievement and representing the highest non-conference year since 2017.

While operating at a deficit, the organization faced some one-time expenses, notably the launch of the registry project and other projects. These decisions, made prudently by our board, lay the groundwork for future growth. Further, we have maintained our reserve account, with six-months operating expenses, and our endowment since they were funded in 2019 and 2012, respectively.

Expenses for the year amounted to just under $250,000, with a strategic focus on scaling up contract positions as we enter 2024. We are confident in our talent retention and scalability model, with a single employee managing a group of independent contractors, enabling us to scale incrementally over the coming years.

Looking ahead, our major commitment is the funding of a portion of expenses for the registry, estimated at around $40,000 annually to be shouldered by Nevus Outreach. This financial obligation aligns with our dedication to advancing research initiatives.

Notably, the "See Beyond" fundraising initiative has been launched—our first ever multi-year fundraising initiative to which our community has already pledged over $100,000. This campaign also incorporates a cadence of friends and family campaigns, a strong monthly giving program and a robust peer-to-peer strategy. These initiatives form a crucial part of our future and enhance the organization’s financial sustainability.

We are proud of the progress made in 2023, especially during a record-breaking Q4 in fundraising since the departure of Morgan Family Foundation funds in 2017. As we enter 2024, Nevus Outreach is poised for continued growth and promise.
AN ELITE GROUP
Outreach Angels give to Nevus Outreach on a regular basis. Their monthly donation provides valuable funding for everything from school resources to medical research. The Outreach Angels are regarded as the most dedicated of all those who give to Nevus Outreach. Their incremental donations transfer across automatically, silently ensuring advance our mission.

JOIN THE MOVEMENT
We are 7 Angels away from our 100–strong Outreach Angel goal! Join hands with us as we soar into the future, wings outstretched. We’ll reach 100, united in purpose, hearts ablaze with compassion. Every hand clasped, every voice raised, fuels our mission. Together, we’ll be the change, the hope, the wings lifting others.

THANKS KATE & COLE FOR YOUR SUPPORT!!
As we step into the new year, we are excited to share with you the remarkable journey and achievements that your support has made possible. The year 2023 was marked by significant strides, including a revitalized brand identity, an interactive website, and numerous initiatives to fortify our community.

Throughout this transformative journey, we’ve come to understand the vast scope of Nevus Outreach beyond its core mission of raising awareness, providing support, and seeking cures for Congenital Melanocytic Nevi (CMN) and related disorders. We have evolved into advocates, cheerleaders, a vital source of information, and a comforting space for connection and understanding.

Recognizing this evolution, our Board and staff collaborated to refine our mission statement, ensuring it authentically represents the inclusive nature of our work. Following thoughtful input from our community, we are thrilled to unveil our updated mission statement:

**Nevus Outreach is dedicated to driving awareness, fostering a supportive community, and advancing research for people affected by congenital melanocytic nevi and related conditions.**

This refined statement encapsulates our rich history while propelling us into a future brimming with possibilities. Our commitment remains unwavering – we will continue to be a steadfast advocate for awareness, disseminating knowledge about CMN globally. We will actively build a vibrant and supportive community, providing a sanctuary for shared experiences, solace, and empowerment. Above all, we will persist in our pursuit of cutting-edge research, pushing the boundaries of knowledge to bring us closer to the ultimate goal of cures.

As we enter this new year, the air is charged with the promise of potential and progress. Unified under our updated mission, we are stronger than ever in our collective endeavor to illuminate the lives of those affected by CMN. Let us stride forward together, hand-in-hand, driving awareness, fostering love, and shaping a brighter future for all.
2023 Year in Review

In 2023, Nevus Outreach became a testament to the extraordinary kindness and momentum within our community, fueled by the passion and contributions of individuals who make a difference. Let’s delve into the impactful moments and the incredible people behind them:

**Branding Transformation:** Kurt Karr and Austin Karr of the MonkeyThis Team, alongside the dedicated branding committee—Chris McKenna, Nate Ruckman, Cristy Phillips, Rita Pink, Jenna Connors, and Wiatt Bowers led the effort.

**Board of Directors Updates:** Merilee Vance, a familiar face returning for a second term on the Board of Directors, shared her invaluable insights after previously serving from 2014-2020. Cara Dolchin, CPA also joined the board after a year of service on the finance committee.

**Representation at AAD Meeting:** Dr. Maggie Mangold and Dr. Diana Olvera brought Nevus Outreach’s spirit to the American Academy of Dermatology meeting in New Orleans.

**Outreach Angels Exclusive Event:** On March 30, 2023, a special gathering allowed contributors to provide feedback on goals for the future of Nevus Outreach.

**Legacy Gift:** The meaningful legacy gift from the Estate of Leland Bradley Stanford, uncle of Kathy Stewart, stands as a testament to the enduring support within our community.

**Congressional Briefing Representation:** Dr. Adrea Bischoff and Lauren Isbell engaged in a dozen meetings with legislators, advocating for funding to the NIH and improved provisions for patient education.
2023 Year in Review

Social Media Takeover Success: Hope Manser, Winnie Dolchin, Fletcher Collins, and Grayson Evans orchestrated a successful social media takeover.

Global Skin’s Elevate Conference Representation: Whitney Casal, our Director of Membership and Metrics, not only represented Nevus Outreach at the conference in Brussels but also actively contributed to launching the GRIDD Study.

New Website Launch: The fresh www.nevus.org, symbolizing growth, opportunity, and vibrancy, became a collaborative achievement, reflecting the spirit of our community.

MyPART Natural History Study: The Nevus Outreach Research Committee, fueled by community input, played a pivotal role in defining priorities for the MyPART Natural History Study, a landmark project conducted fully with federal interest.

Forward Together Virtual Conference: Hosting this event online allowed us to connect, share research findings, and announce St. Louis as the conference site for 2024. Special thanks to Whitney Casal and Lilly Hennesey for co-hosting.

Rare KC Event at Children’s Mercy Research Institute: Lauren Isbell represented Nevus Outreach learned about the latest research and clinical care for neurological rare diseases. After a full day of presentations, the Budine Family the Sturgeon family at Waldo Pizza.

Board Retreat and Friends and Family Campaign Launch: During a two-day retreat in St. Louis, the group planned for the organization’s future and launched their personal Friends and Family Campaigns, raising nearly $20,000.

PeDRA Conference: Dr. Maggie Mangold and Lauren Isbell, representing Nevus Outreach in Atlanta at the PeDRA Conference, formalized the next steps for the NOOR (Nevus Owners Outcomes Registry) set to open for enrollments in Summer 2024.

End of Year Success: As 2023 concluded, the community’s support propelled Nevus Outreach to surpass revenue goals, achieving the highest non-conference year since 2015. The top ten personal
Donors—Donna Sue Rosene, Erik Gregory, Joanna Pozen, Robert Brown, Cristy Phillips, Beverly Rizer, Maggie Mangold, Judith Goldberg, Diane French, and Steve Bischoff—played a pivotal role in this success. On December 31 with a full-circle moment in mind, Nevus Outreach founder Jeff Curtis called in a last-minute donation to be our closer on New Year’s Eve to bring our total revenue for the year to $210,818—a number that put us at 112% of our stretch revenue goal for the year signaling the highest revenue since the Morgan Family Foundation departure.

In 2023, Nevus Outreach thrived, thanks to the personal connections, dedication, and support of our community members, partners, and contributors who truly make a difference.

THANK YOU
TO THE BOARD OF DIRECTORS FOR LEADING THE CHARGE FOR THIS MILESTONE YEAR.

WIATT BOWERS - CHAIR
LISA O'BRYAN - VICE CHAIR
OLIVER HOLMES - SECRETARY
CARA DOLCHIN - TREASURER
MAGGIE MANGOLD
CRISTY PHILLIPS
MATTHEW ROGERS
ANDREA BISCHOFF
ERIK GREGORY
MERILEE VANCE
JENNA CONNORS - PAST BOARD CHAIR
Because of your generosity Lauren represented Nevus Owners and families at the Pediatric Dermatology Research Alliance (PeDRA) Corporate Council Meeting and at a meeting with the National Institutes of Health at their headquarters in Bethesda, Maryland. These events were invitation-only and Lauren was selected as one of three participants nationally to bring the voice of the patient experience to the table.

With this partnership, we are able to extend our research efforts further. Through our partnership with PeDRA, Nevus Outreach has the opportunity to:

- Connect with other researchers worldwide who are working on similar projects
- Share our research findings and learn from the experiences of others.
- Get involved in the development of new research initiatives.
- Help to shape the future of research
- Show commitment to our mission of advancing research for CMN and related disorders.

Remember that time we all gathered for the 2008 Conference? Or Freewheel in 2014? The laughter, the support, the feeling of belonging like never before?

We want to take a trip down memory lane with you!

Share one of your favorite moments, events, meet-ups, or fundraisers at [nevus.org/fta](http://nevus.org/fta)

Don’t forget to include a picture if you have one! We love seeing your smiling faces and the moments that have shaped our Nevus journey.

Here’s to cherishing the past and embracing the amazing future that awaits us!

SHARE A BLAST FROM THE PAST
NEVUS.ORG/FTA TODAY

THANK YOU

Pediatric Dermatology Research Alliance

Nevus fam, let’s take a trip down memory lane!
PhRMA Foundation

Nevus Outreach is delighted to announce that we have been honored with a $3,000 grant from the PHRMA Foundation. This grant is a testament to the impactful work we do in supporting individuals and families affected by congenital melanocytic nevi. The PHRMA Foundation, dedicated to advancing public health through research and education, recognizes the importance of our mission in the dermatological community.

With this generous grant, Nevus Outreach plans to further expand its outreach programs, raise awareness about nevus-related conditions, and provide crucial support to those navigating the challenges associated with congenital melanocytic nevi. The support from the PHRMA Foundation reaffirms the value of our initiatives in promoting dermatological health and well-being.

We extend our sincere gratitude to the PHRMA Foundation for their commitment to advancing public health and their recognition of the significance of our work. This grant will undoubtedly have a meaningful impact on our ability to reach and support more individuals affected by congenital melanocytic nevi.

For more information about the PHRMA Foundation and their commitment to improving health through research and education, please visit: https://www.phrmafoundation.org/

Thank you once again to the PHRMA Foundation for their support and belief in the mission of Nevus Outreach.
As we step into the new year, we’re thrilled to share the exciting vision we’ve set for our membership program in 2024. These objectives aim to empower, excite, and most of all better serve our beloved community. Experience the growing excitement as we take our membership program to new heights!

**Goal 1: Expand Member Resources**

This year we’ll be exploring options for a private, online space available exclusively to our members. We imagine a virtual portal where you can access all kinds of valuable resources and even connect with other members (no Facebook required!). In addition, we aim to develop a treasure trove of recommendations straight from the lived experiences of our community members. After all, who better to guide and inspire than those who’ve walked a similar path? Ultimately, we want to provide a better platform for the voices that matter most: yours.

**Goal 2: Develop Community Support**

Strengthening our community is at the core of our mission. We also recognize the importance of sustaining connections beyond the in-person conference that takes place every two years. In 2024 and beyond, expect increased opportunities for member interaction. Whether it’s regional in-person gatherings, virtual meetups, themed discussions, or collaborative projects, we’re committed to keeping the conversation alive and vibrant.

As a step towards growing and sustaining our efforts around support, Nevus Outreach is committed to the establishment of a Support Committee in 2024. Led by staff and supported by Board Members, Members’ needs will be at the forefront of these activities. One specific goal from this committee is to develop a structure for ongoing support groups, tailored to specific needs and experiences. Everyone’s journey is unique, and we want to ensure you have the support you need, when you need it. Whether you’re navigating a specific challenge or celebrating a triumph, our community will be here for you every step of the way.

As for our international members, we’ve heard your call! Our goal is to make resources more accessible on a global scale. This means not only broadening the reach of our content but also incorporating different languages to ensure that every member, regardless of their location, feels at home in our community.
Join the Excitement

But we’re not just telling you about these incredible changes – we want you to be an active part of them! If you haven’t already, we invite you to become a member of Nevus Outreach today. Our free membership opens the door to a world of resources and connections. For those seeking a bit more, our supporting membership (available for just a small yearly fee) comes with added benefits, ensuring an even richer and more personalized journey.

As we embark on this ever-growing adventure together, we invite you to join our community at Nevus Outreach. Your membership not only grants you access to exclusive features but also strengthens the bonds of our incredible community, and supports the growth of Nevus Outreach as an organization. Together, let’s make 2024 a year of stronger connections, unwavering support, and shared success!

YOU BELONG HERE

At Nevus Outreach, we understand the power of community. As a member, you support our mission of creating a robust network for advocacy, support, and research. Help us revolutionize the way our loved ones live with their rare condition, and create a brighter future for people with congenital melanocytic nevi (CMN)!

or visit: nevus.org/membership
Beyond Cash

How Non-Cash Donations Fuel Nevus Outreach Efforts

While financial contributions are vital for Nevus Outreach, there’s an often-undervalued way to make a significant impact: donating non-cash assets. It’s a win-win situation, allowing you to potentially gain tax benefits, and support this crucial cause — all without dipping into your cash reserves.

What are some qualifying assets?

- Stocks and mutual funds: Donate appreciated securities for fair market value and avoid capital gains tax.
- Charitable life insurance: designate Nevus Outreach as a beneficiary.
- Donor Advised Funds: We are eligible to receive gifts from your DAF.
- Give directly from your IRA to satisfy your Required Minimum Distributions starting at age 73.

Why choose non-cash donations?

Maximize your impact: Donating appreciated assets often translates to more resources for Nevus Outreach compared to cash equivalents.

Potential tax benefits: Depending on the type of donation and your tax situation, you may be eligible for tax deductions.

Ready to get started?

Nevus Outreach welcomes non-cash donations and can guide you through the process. Here are some steps:

1. Contact Nevus Outreach: Discuss your donation goals and explore accepted items.
2. Consider your options: Research non-cash donation platforms and choose the method that best suits your asset.
3. Spread the word: Encourage others to explore non-cash giving as a powerful way to support Nevus Outreach.

Remember, your non-cash has the potential to make a real difference. By thinking beyond cash, you can unlock a hidden wellspring of support and fuel the critical work of Nevus Outreach.

When you are ready contact us at: info@nevus.org

*Stock initially purchased at $2,000

Sell First, then Give Sells for $20,000

Give First, then Sell Sells for $20,000

Gift to Nevus Outreach $15,716

Gift to Nevus Outreach $20,000

Capital gains tax 20%, NII 3.8% $4,284

NO capital gains tax on charitable donations of stock

$0
We are so thankful for Nevus Outreach.

Without them, we would still be scared, confused, and badly misinformed. Instead we are educated and happy that Aiden never has to feel alone with his condition.

—Michael & Melissa

**HOW TO GET INVOLVED WITH NEVUS OUTREACH**

Nevus Outreach is always looking for a diverse set of talents to help us move our mission forward. Our organization is presently organized with the following committees to carry out our work:

- **Advocacy**: Drives policy change to ensure access to quality care research advancements via partner organizations.
- **Awareness**: Reaches our target audience, builds awareness of CMN and our mission and mobilizes broad support.
- **Conference & Events**: Ensures events create a supportive and inclusive environment.
- **Finance**: Paves the way for Nevus Outreach to effectively manage our resources and achieve our mission.
- **Fundraising**: Provides the financial resources needed to carry out our mission. A successful fundraising strategy requires a clear understanding of the our goals, a well-planned approach, and effective communication with donors and stakeholders.
- **Research**: Identifies key issues and opportunities to advance research priorities.

**Research Committee Working Groups**: is looking to launch three working groups focused on: Education of NICU & OBGYN staff; Developing Opportunities for Medical Students; and Navigating life with CMN—development of educational resources and addressing psychosocial aspects of the condition.

**Support & Wellness**: Launching 2024 with the hope of fostering a supportive community through various support groups focusing on mental health and emotional well-being.

**Chapters Exploration Committee**: Launching Spring 2024 Committee will analyze the “regional group” legacy program and evaluate for next steps with the possible evolution to a local chapter model.

To learn about committee service, reach out to Whitney at whitneycasal@nevus.org
We are thrilled to share a significant development in the journey of Nevus Outreach—the execution of a crucial contract by our Board of Directors with the Pediatric Dermatology Research Alliance (PeDRA). This strategic partnership marks a pivotal step towards the imminent launch of our highly anticipated patient registry, the Nevus Owners Outcomes Registry (NOOR).

The collaboration with PeDRA holds immense significance as it enables the co-creation of a patient registry adhering to the best practices established by the National Center for Advancing Translational Sciences (NCATS). This joint effort mitigates the challenges associated with aligning with a singular academic institution, ensuring a comprehensive and inclusive approach to our registry.

We are very grateful to Dr. Harper Price, Dr. Carrie C. Coughlin and others and their respective institutions, as well as PeDRA for their collaborative efforts on the project which will establish a secure and research-ready database. This partnership aligns with our dedication to advancing nevus-related research while navigating the complexities of regulatory standards.

Dr. Maggie Mangold, who led our research committee from 2021 to 2023, joined PeDRA staff earlier this month to assume oversight of this groundbreaking project. With the collective expertise of Nevus Outreach and PeDRA, we anticipate securing the necessary approvals and commencing patient enrollment by summer 2024.

This partnership signifies not only a collaboration but also a strategic commitment to elevating the future of our community through cutting-edge research and innovation. Stay tuned for further updates on this momentous initiative.
Mark your calendars, Nevus FAM! This isn’t just a conference, it’s a community celebration where nevus owners, loved ones, and experts collide for three days of epic knowledge, heart-warming connections, and enough fun for a lifetime of memories!

Sunday evening sets the stage for epic bonding with a casual Meet & Greet. Mingle with old and new friends, share stories, and let the love flow.

Monday morning brings a power-packed kickoff with updates on Nevus Outreach’s amazing work, the lowdown on Nevus 101, and the unveiling of the game-changing Nevus Outreach Registry Project! While you’re soaking up the knowledge, your littles and teens will be busy with hotel activities and a trip to Slick City, the ultimate slide paradise! And who can forget the legendary Pizza & Pool Party that night? Get ready for cheesy slices, splashing good times, and memories that’ll last a lifetime.

Tuesday’s a brain-boosting bonanza with insights from top experts in dermatology, neurology, and even plastic surgery! You’ll learn everything from conquering your nevus like a pro to unlocking its unique mysteries. Meanwhile, your mini-mé’s will be conquering an off-site fieldtrip!

As the day winds down, enjoy some jaw-dropping entertainment and a moving memorial service to remember our nevus friends gone too soon.

Wednesday’s your day to shine even brighter! Adults will dive into navigating life with a nevus, fueled by the inspiring stories of fellow nevus owners. And to cap off this incredible week, get ready for our silent and live auctions bonanza, dinner and dance party so epic it’ll make the disco ball jealous!

But the fun doesn’t stop there! Your grand finale is a trip to the world-famous St. Louis Zoo, where you can bond with your nevus fam outside the conference walls. This isn’t just a conference, it’s a celebration of your strength, your community, and your unique, beautiful nevus. It’s a chance to shine brighter than ever, learn, laugh, and make memories that’ll stay with you long after you leave St. Louis.

So grab your friends, your family, and your brightest smile. We’re waiting for you in St. Louis, July 7-11! Let’s SHINE together!
FUNDRAISE YOUR WAY TO CONFERENCE

Need help getting to conference? Now worries we got you!
Hosting a Grassroots Fundraiser can help you with
REGISTRATION and more

Get ready to make a real impact on the Nevus Outreach Conference with our awesome toolkit designed for grassroots fundraising. This toolkit is like your go-to guide for anyone eager to champion the cause and reduce their own conference costs.

We’ve got creative and effective fundraising ideas tailored to fit everyone’s style, from social networks to local hangouts and online platforms. Engaging in grassroots initiatives isn’t just about supporting the conference—it’s your chance to shape your own experience and be part of a community that’s got your back.

As funds roll in through grassroots fundraising, we’re making sure the conference stays inclusive and affordable for all. Collaborate with others, pool your resources, and let’s make the July 7–11, 2024 conference in St. Louis, MO, a success together. The toolkit even gives you tips on storytelling and communication to help you spread the word about why grassroots support is so important. So, get ready to dive in, make a difference, and be a key player in making the conference accessible to everyone!

Greatful to recognize the following people for hosting fundraisers:

Brucklacher Family and the Liberty VFW Post 6755
Blood by Fire Motorcycle Club in honor of Abigail Clark and the Graeff family
Friends of Georgetown Basketball
Clara Barton Elementary School
Tyler Place Tribute to the Connors, Organized by the Mangold Family
Connor’s Newburyport Cruise

SHOUT OUT
Download the grassroots fundraising toolkit here!

or visit: Nevus.org/grassroots-fundraising

Check out more photos and tips for great events here!

or visit: NevusConference.org/wakelet

August Holmes’ Fifth Birthday
Angel Bermudez Birthday
Evan Coleman’s Angelversary
Rylee O’Bryan College Graduation
Katie Vann and Brownings United Methodist Church
Ant’s Army and the Vanderzee Family
Kristen Talbot Flipside Watersports Cable Freeze Event
Nevus Outreach, in collaboration with Global Skin, is thrilled to introduce Rally for Rare—a nationwide initiative hosting 11 events in 11 days across the United States! From February 27 to March 9, 2024, these gatherings will focus on fostering connections among nevus families. We’re excited to announce the cities where these impactful events will take place:

- SAN DIEGO, CA
- DENVER, CO
- AUSTIN, TX
- KANSAS CITY, MO
- COLUMBUS, OH
- PHILADELPHIA, PA
- WASHINGTON D.C.
- RALEIGH, NC
- JACKSONVILLE, FL
- NEW YORK, NY

Rally for Rare is more than a series of events; it’s a heartfelt initiative to bring our community together, providing support, camaraderie, and a platform to share experiences. Join us on this incredible journey as we connect nevus families and make lasting memories.

Stay tuned for further details and updates—it’s shaping up to be an unforgettable experience!
Rare Disease Day holds profound significance for the Nevus community, providing a unique opportunity to shed light on the challenges and triumphs of those affected by congenital melanocytic nevi (CMN) and related disorders.

This day is a powerful catalyst for raising awareness, fostering understanding, and garnering support for individuals with rare conditions. Nevus Outreach is thrilled to announce an all-day online fundraising blitz, reminiscent of the success of Giving Tuesday 2023.

Join us on February 29th as we come together online for a day filled with engagement, shared stories, and collective efforts to make a tangible impact. Mark your calendars, because on this special day, all funds raised will be directed to PeDRA (Pediatric Dermatology Research Alliance) to further support the NOOR (Nevus Owners Outcomes Registry).

Your participation will play a crucial role in advancing research and improving outcomes for those affected by CMN. Let’s unite on Rare Disease Day 2024, contribute to the cause, and make a lasting difference in the lives of the Nevus community.
Nevus Outreach would like to extend a heartfelt invitation to the 3rd CSD Hill Day Event, taking place from April 21 to April 23, 2024. As we gather for this impactful occasion, we have the opportunity to connect, advocate, and make a lasting difference in the lives of dermatology patients.

The Coalition of Skin Diseases has a long standing history of partnership with Nevus Outreach. Our executive director Lauren Isbell, is the current Secretary of the CSD Board of Directors, is enthusiastic about this event.

Andrea Bischoff, a dedicated Board Member who attended the CSD Spring Hill Day in 2023, shared, "Participating in Hill Day was an empowering experience that allowed us to voice the needs of our community directly to legislators. It’s a unique chance to make a real impact."

The event includes a welcome reception on April 21, in-person training, a social event/dinner on April 22, and meetings with legislators alongside a congressional briefing/lunch on April 23.

To make this event accessible, the CSD is inviting an additional 15 local advocates to join our efforts. The application period is open until March 1, with our Advocacy Task Force reviewing and informing applicants of their status by March 4.

We encourage representatives from all our groups to join this unique opportunity to connect with peers and advocate for the dermatology patient community.

If you are unable to attend, please share this invitation with those in your patient population whom you believe would be a good fit.

Don’t miss this chance to be part of a meaningful advocacy initiative. Apply here and be a crucial voice for the Nevus community.

Hope to see you there!

SAVE THE DATE

ARE YOU IN THE WASHINGTON, D.C. AREA?

or visit: Nevus.org/hillday
HELP FUND THE FUTURE OF SUPPORT, AWARENESS, AND RESEARCH.

Consider this your official invitation to become an active participant in creating positive change for the Nevus community.

By pledging support, hosting fundraisers, making introductions, joining the Outreach Angel program, or including Nevus Outreach in estate plans, you can contribute to our success and help transform the lives of those affected by congenital melanocytic nevi (CMN) and related conditions.

SCAN HERE TO SHOW YOUR SUPPORT
or visit: Nevus.org/see-beyond

Yes! I would like to support the Nevus Outreach with my gift...

☐ $5,000  ☐ $1,000  ☐ $500  ☐ $250  ☐ $75  ☐ $25  ☐ Other
☐ I’d like to make a recurring gift of $_____ per month.
  ☐ Please contact me if I would like to host a fundraiser in my community.
  ☐ Please contact me if I would like to introduce Nevus Outreach to someone.
  ☐ Enclosed is my check made payable to Nevus Outreach.
  ☐ Charge my donation to my: ☐ Master Card  ☐ Visa  ☐ Discover  ☐ Am Ex.

Mail your gift to Nevus Outreach 361 Southwest Drive #353, Jonesboro, AR 72401

Name as it appears on card __________________________________________

Account Number_________________________ Expiration Date ________ CVV

Signature ___________________________________________ Date __________

☐ Contact me about making a provision for Nevus Outreach in my will. __________

☐ My employer will match my gift. The matching form is ☐ enclosed ☐ will follow.

Name___________________________________________________________

Address__________________________________________________________

City_________________________ State__________ Zip Code____________

Phone ___________________________ E-mail Address _____________________

My gift is in honor of/in memory of ________________________________

Please notify the following recipient: Name__________________________________

Address___________________________________________________________

City_________________________ State__________ Zip Code____________

Nevus Outreach is a 501(c)(3) Charitable organization and contributions are tax deductible to the extent allowable by law. EIN: 59-3455128