



June 24, 2021
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STEP for Bleeding Disorders 5K Walk/Run Fundraiser
November 6, 2021
Saluda Shoals Park
6071 St. Andrews Rd
Columbia, SC 29212

Dear Friends, Supporters and Sponsors:

Imagine receiving news about your health or the health of a loved one that will impact you for the rest of your life. You have a bleeding disorder. Your career, hobbies, where you live, family planning, and many other life decisions are now all affected. Your doctor can prescribe a treat regimen, but where do you turn for help and support in between office visits. Who has the most relevant experience or practical knowledge to help you navigate life with a rare and life threatening chronic disorder?

Being part of a community that shares experiential knowledge often means the difference between effectively managing a chronic disorder like Hemophilia or von Willebrand Disease, or being debilitated by it. Patient education, public awareness, support services, and advocacy for those affected are the principals at the core of our organization. We are *The Bleeding Disorders Association of South Carolina (BDASC)*, and we would love to have your support!

As South Carolina's leading resource for children and families affected by bleeding disorders, the BDASC maintains close relationships with our community, providing them the opportunity to network with other families experiencing the same situations, seeking guidance and emotional support for an often anxiety-ridden and frightening diagnosis. We support our families from the initial diagnosis throughout their lifetime. We serve the entire state of South Carolina, over 1,000 individuals and their families, and we are a tight-knit community family. Hemophilia and other bleeding disorders are rare and the occurrence of knowing someone in your lifetime with the same disorder is highly unlikely. BDASC helps our members develop a strong support system that will benefit the emotional health of individuals within the family unit. Having a bleeding disorder can be an extremely isolating experience and the feelings of knowing you are not alone in this chronic health struggle are unmatched in importance.

Bleeding Disorders Association of South Carolina is a 501(c) 3 non-profit organization which serves the state of South Carolina and is a chapter of the *National Hemophilia Foundation (NHF)* and a member organization of *Hemophilia Federation of America (HFA)*. BDASC is proud to present our **"STEP for Bleeding Disorders 5K Walk/Run Fundraiser"**. The event will be held at the Saluda Shoals Park in Columbia, South Carolina, on November 6, 2021. This event is our Chapter's largest fundraising event and is made possible by the generous support of our donors and sponsors. The funds raised will support the educational programs, advocacy initiatives, and supportive services we provide to those affected by bleeding disorders in South Carolina. A small portion of the walk proceeds will be provided to the research of the National Hemophilia Foundation. Walks such as ours occur each year throughout the nation to support local chapters.

Persons with bleeding disorders, like hemophilia, have clotting factors or proteins in their blood that are missing, low or do not work as they should. The lack of a specific clotting factor prevents the blood from clotting normally. This can lead to bleeding into joints, muscles and internal organs. Bleeding into the joints and muscles is extremely painful and causes long term damage. Certain internal bleeding episodes such as bleeding in the brain, can be life-threatening. Treatment is expensive and can be complicated. Currently, there is no cure.



For the past 48 years, **Bleeding Disorders Association of South Carolina**, (formerly know as Hemophilia of South Carolina) has provided support services and educational programs to the South Carolina bleeding disorders community. Below are just some of the programs and services that your sponsorship dollars will support and ensure to continue. Our advocacy efforts have improved the lives of many in our community.

We provide the following support services, free of charge:

1. An Annual *State-Wide Annual Meeting and Educational Conference*
2. *Support groups, workshops*
3. *Camping support* to our state camps for children affected by blood disorders
4. *College Scholarships*
5. *Virtual Technology Managed Devices Programming (VIP)*
6. *Financial Assistance Program*
7. *Patient Advocacy Training*
8. A state legislative day in *March, the National Bleeding Disorders Awareness Month*, to raise awareness of bleeding disorders and advocate for the needs of those affected.
9. Advocacy *Coalition* and Ambassador Program
10. Annual *teen & young adult support/educational events*
11. *Walk and Golf Fundraiser* to raise awareness and funds
12. *An outreach and referral program* in collaboration with our Hemophilia Treatment Centers
13. Annual year-end winter meeting and *Holiday* gathering
14. *Quarterly newsletter* for our members, all affiliations of our organization, state hospitals, state agencies and HTC's.
15. *Educational and social dinner programs and events* throughout the state to assist families in their *local areas*
16. *Educational Family Camp and Adult Retreat Weekend*
17. *Local, regional and national training* and travel sponsorships
18. *Participation at Washington Days at the Nation's Capital.*
19. *Funding* to support the *NHF Judith Graham Pool Research* Fellowship and funding to *world projects* for underdeveloped countries for treatment.
20. *BDASC sponsors children around the country with financial aid through the Save-One-Life Organization.*

We hope we can count on you to support our mission. Hundreds of families in South Carolina will benefit from your support and on behalf of **Bleeding Disorders of South Carolina**, we thank you for your consideration. Should you have any questions, please don't hesitate to call or e-mail me at the below contact information. Additional information on the walk, participation and sponsorships packages can be located on our website at www.hemophiliasc.org.

Warmest Regards,

Suzanne Martin, Executive Director
Sue.martin@hemophiliasc.org
(864) 350-9941