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Fall 2021 #2 Newsletter: National Neutropenia Network



Join us for our Virtual Event!

'Steps for Neutropenia' is a virtual race to connect your family and friends across the world to help raise awareness about neutropenia and raise funds to keep it strong. Your contributions will enable us to provide education, support, and connections to experts for children and adults with chronic neutropenia.

This virtual event takes place anywhere and takes place from October 1 through October 31, 2021.

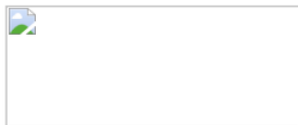
Set a target to reach 10,000 steps each day! You can run, walk, swim, bike, dance, knit, read-create your own path to support those with neutropenia and help the network stay strong! Teams can be formed as well! You can also connect it to your Fitbit if you have one!

Registration is open and event continues through October 2021.

[You can find registration and information through our website.](#)

Join the fun, raise awareness, and earn those steps!

Thank you for supporting the National Neutropenia Network.



Play Groups for Kids, ages 5-18.

The Barth Syndrome Foundation is hosting 'Play Groups' for kids through 'Teleplay' where kids with rare conditions are invited, ages 5-13; however they do have breakout rooms for kids up to age 18. Parents will need to register their children in advance.

It starts at 5:00 PM EDT. The group meets every other Thursday and is led by medical students. The next gathering is this Thursday, Oct. 7th at 5:00 PM EDT. To register: <https://sforce.co/3i48fd8>.

Helpful Resources

Emergency Room Card
[English ER card \(PDF\)](#)
[Español ER tarjeta \(PDF\)](#)

[ANC Calculator](#)

[E-Cards: submit & send](#)

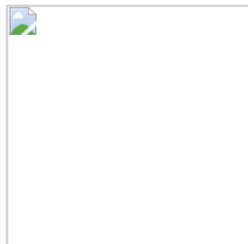
Information on types of neutropenia, management, and more at:

[The Severe Chronic Neutropenia International Registry](#)

[Ella Jewel Foundation](#)
[Cohen Syndrome Association](#)
[Barth Syndrome Foundation](#)

Neutropenia Warrior Shirts

Jamie has created a new shirt fundraiser that benefits the National Neutropenia Network. Thank you, Jamie!



[Click here to check them out.](#)

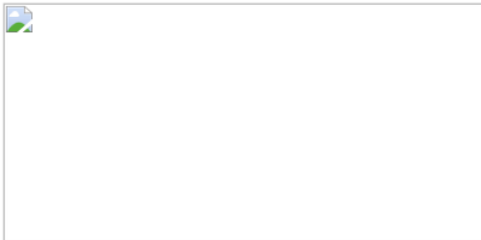
**Sales end soon!
Happy Shopping!**

Genetic Testing Webinar

Learn more about Path4ward and Genetic testing.

[Genetic Testing Webinar](#) Access
Passcode: C#c0k6=E;

Recorded September 10, 2021, in collaboration with Dan Link, MD, Precision, Genome Medical, and X4



Twenty-seven years, on October 26, the State of Michigan incorporated the National Neurotopenia Network as a non-profit organization!

Happy Birthday, NNN!

Thanks for Subscribing

The team at [National Neurotopenia Network](#) wants to extend a hearty 'thank you' to all of our wonderful readers. You've helped make our community shine, and we want to keep that going! For more information on our mission and how you can help visit [our website](#).

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Give Today

The National Neurotopenia Network depends on support from the friends and families of those whose lives have been forever changed and challenged by this rare disease. Thank you for your support.