Can't Join Us In Person? Run a Virtual 5K!

What exactly IS a Virtual 5K?
It is just like any other 5K, except the race location is wherever you are! You get to choose where, with whom, and even how you want to complete the 5K. By participating virtually, you can run the 5K at any time leading up to March 26th. It can even be done in increments - keep track of your distance until you hit the 5K.

Help Us Raise Funds and Awareness
This event is open to individuals of all ages and abilities. Once you register, either as an individual or as part of a team, you are set. We would love to see the posts of pictures participants on their 5K route. This is completely optional - but since our goal is to bring awareness to MPS and ML, we want to do just that. And by using #RaleighRun2022 and #MPSAwareness on your social media posts, we will be able to keep up with the fun!

Register, Sponsorships & Donations

- **Platinum Sponsor:** $1,500
  Includes 25 participants and shirts, recognition on shirts and website recognition. Name on banner at the event.

- **Gold Sponsor:** $1,000
  Includes 25 participants and shirts, recognition on shirts, website recognition. Small sign at the event.

- **Team Sponsorship:** $500 and $300
  $500: Includes 25 participants and shirts, recognition on shirts and website recognition.
  $300: Includes 15 participants and shirts, Recognition on shirts, Event, and Website Recognition.

- **Individual 5K Runner:** $25
  Includes shirt

- **1 Mile Walker:** $15
  Includes shirt

To Register Online, Visit: bit.ly/runforrare22

Event Contacts:
Madeline Schehl, madeline@mpssociety.org
Tracy Kirby, tracy@mpssociety.org

The Mission of the National MPS Society

*Who We Are:*
The National MPS Society exists to cure, support and advocate for MPS and ML. We provide hope and support for affected individuals and their families through research, advocacy and awareness of these devastating diseases.

Since 1974 the Society has supported individuals and families affected by MPS and ML. We are governed by a member-elected volunteer Board of Directors, many of whom are parents of children with MPS. We also benefit from the expertise of a Scientific Advisory Board, comprised of world-class physicians, researchers and medical professionals throughout the world.

*How We Help:*
The Society is the national support for families and children affected by MPS and ML. The Society advocates through efforts in Research, Family Assistance, Legislative Advocacy, Educational and Fundraising Committees. The Society has funded over $11 million in research.

*How You Can Help:*
We need your support to help us educate others about our mission and to raise the money we need to support medical research — the key to longer, happier lives with MPS and ML.
Raising Funds for Rare Genetic Diseases

What is MPS and ML?
Mucolipidosis (ML) and Mucopolysaccharidoses (MPS) are genetic lysosomal storage diseases (LSD) caused by the body’s inability to produce specific enzymes. Normally, the body uses enzymes to break down and recycle materials in cells. In individuals with MPS or ML the missing or insufficient enzyme prevents the proper recycling process, resulting in the storage of materials in virtually every cell of the body.

To learn more about MPS, ML, and the Society, visit: mpssociety.org

Race Course
5K Run and 1 Mile Walk
Awards for 8 tiers of mens/womens finishers

Use this QR Code to visit
The Raleigh Run Website:

National MPS Society
March 26 at 8:00 AM
Dorothea Dix Park
Website: bit.ly/runforrare22
#RaleighRun2022 #MPSAwareness