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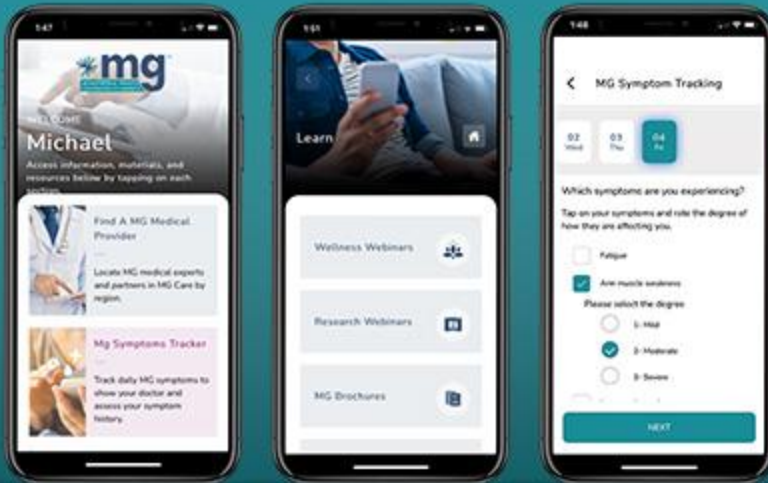
*Brought to you by the Myasthenia Gravis Foundation of America*

APRIL 2022

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FEATURED

**MyMG  
Mobile App**



## Introducing the Brand New MyMG Mobile App – Now Available for Download

MGFA has already added new features and improved the MyMG Mobile App since we launched 2 weeks ago. Get it on iPhone or Google Play platforms and manage your myasthenia gravis in a new and personal way - from the palm of your hand.

[LEARN MORE](#)



Talk about generalized myasthenia gravis (gMG) with another patient or caregiver

[ENROLL NOW](#)

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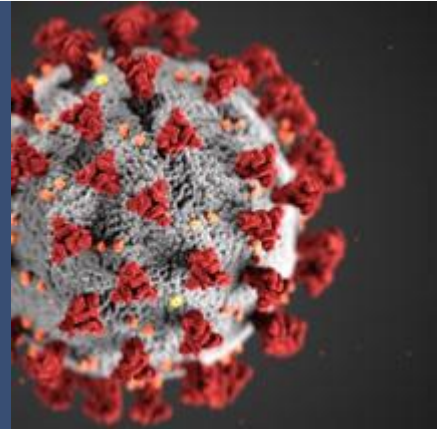


## Janet Myder Was the Epitome of Dedication to MG Advocacy

Janet Myder, a truly well-known friend of the MG Community, passed away earlier in April. We honor Janet and recognize her many volunteer contributions and the influence she had on all of us. [Learn More»](#)

## New Research: Safety and Tolerability of COVID Vaccination in MG Patients

Our data supports the safety and tolerability of mRNA-COVID-19 vaccines, which should be strongly recommended in MG patients who could be at higher risk of complications if exposed to SARS-CoV-2 infection. [Learn More»](#)



**VYVGART™**  
(efgartigimod alfa-fcab)

New FDA-approved treatment  
Talk to your neurologist  
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### Preparation Before You Have an MG Crisis



### When Rare Disease Advocacy is Personal



### Identifying MG Weakness – A Patient Perspective

You should have materials and information available BEFORE an MG crisis occurs so you or your caregiver can act fast. Here are tips to prepare.

[LEARN MORE](#)

Four years ago, Brenda was living in Miami and working in marketing. Now, she is involved in rare disease advocacy. It's personal because she has MG.

[LEARN MORE](#)

“Weakness – it was never in my vocabulary until I was diagnosed with MG. Maybe that’s why it took me 26 years to be diagnosed.”

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## Start Taking Action Now for MG Awareness Month in June

We are Turning Awareness into Action again this June to find new and creative ways to make sure people around the world learn more about MG. Start taking action right now by asking for proclamations, working with the media, sharing videos, and planning events. [Learn More»](#)

## Apply to Participate in MG Clinical Trials

MGFA maintains a list of MG-centric clinical research trials. Check out the list, and consider applying to participate as a patient in trials near you. [Learn More»](#)



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## Gail Duckworth Memorial Garden

*Honor a loved one with a donation to our  
new MGFA virtual memorial garden.*

MAKE A MEMORIAL GIFT

CHALLENGE

## Join the MGFA Coast-to-Coast 2740 Challenge

*Create a team, or join one, and start  
fundraising today to support our work.*

SIGN UP TODAY

Is this monthly e-newsletter valuable for you and your family? Let us know. You can send comments directly to [MGFA@myasthenia.org](mailto:MGFA@myasthenia.org) or contact Michael Antonellis at [mantonellis@myasthenia.org](mailto:mantonellis@myasthenia.org).



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