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DECEMBER 2021

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FEATURED



FDA Approves New Myasthenia Gravis Treatment

The FDA has approved a new treatment for MG patients called VYVGART (formerly efgartigimod). It was developed by argenx with support from the MGFA and our Medical and Scientific Advisory Board.

[LEARN MORE](#)

See if you can help inform myasthenia gravis research
The more that's known, the more that's possible.

MyRealWorld™ MG
[JOIN THE STUDY*](#)
*Eligibility requirements apply.

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REGISTRATION OPEN for the



Annual MGFA National Patient Conference

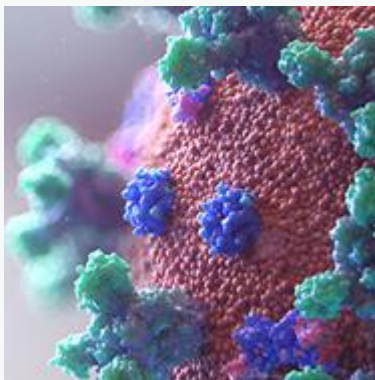
Our annual MGFA National Patient Conference will again be virtual. This exciting experience brings together MG patients, caregivers, partners, and medical professionals. [Register»](#)

Today is a Good Day to Give

Our annual year-end appeal is in full swing, so please consider supporting new MG research funding and programs that help the MG Community. Today is a Good Day to Give in support of the work that MGFA is doing every day. [Donate»](#)



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Special Report: COVID-19 Vaccine Survey Data Presented



MGFA Board of Directors Appointed to New Positions

Congratulations to
Brian Gladden, Bill



International Conference Scheduled for May 2022

During MGFA
International
Conference, global

MGFA Medical professionals presented COVID-19 vaccination data during December 6th Research Webinar.

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Sauerwine, and Celia Meyer for accepting new leadership positions on the MGFA Executive Board.

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myasthenia experts come together to present progress against MG. We meet in May 2022 – abstracts due before January 17.

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Inspired by **patients**.
Driven by **science**.

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OUR MG VOICE

**Registration open for
Rare Disease Week on
Capitol Hill starting in
February**

[LEARN MORE & SIGN UP](#)

RARE
ACROSS AMERICA

EVERY VOICE. IN EVERY DISTRICT. MATTERS

The logo features a stylized sun with rays in orange and yellow, positioned above the text 'EVERYLIFE FOUNDATION FOR RARE DISEASES'.
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ALEXION[®]
AstraZeneca Rare Disease

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VOLUNTEER

MGFA Offers So Many Ways To Get Involved

Take Action Today!

LEARN MORE

Research Webinar: Adverse Event Unit Project

View our latest webinar featuring Dr. Michael Hehir focused on the increasing emphasis of side effect burden for patients with myasthenia gravis. [More»](#)



EVENTS

Start a Community Event Where You Live

We can help!

LEARN MORE



Descartes-08

A new Phase I/II investigational RNA cell therapy
for patients with generalized Myasthenia Gravis

NCT04146051

Learn more

Cartesian
THERAPEUTICS

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Happy Holidays and Here's to a Happy, Healthy, and Successful New Year

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



Myasthenia Gravis Foundation of America
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