Who We Are

Meet our Board of Directors

Dr. Leslie Cooper, MD – Chair of the Cardiovascular Department at the Mayo Clinic in Jacksonville, Florida

DeLisa Fairweather, PhD – Associate Professor and myocarditis researcher, Director of Translational Research, Department of Cardiovascular Medicine at the Mayo Clinic in Jacksonville, Florida

Dr. Jack Price, MD – Associate Professor of Pediatrics at Baylor College of Medicine and the Clinical Director of the Cardiovascular Intensive Care Unit at Texas Children’s Hospital

Candace Moose – Giant Cell Myocarditis survivor

Joseph Rumore – Myocarditis survivor and former Managing Director of a national insurance company

Louis Romano – Owner of Home Well Senior Care, a home health care agency

Joel Aranson – Founder and Chairman of National Sporting Good Corporation and father to a Myocarditis victim

Randy Vanness – Community leader and father to a Myocarditis victim

Christopher Corso – Vice President and Senior Reinsurance Placement Officer XL Catlin Insurance and father to a Myocarditis survivor

Francine Andrea – Director – Vice-President for Enrollment Management, Student Affairs and Chief Compliance Officer for Felician University

How You Can Help

The Myocarditis Foundation is seeking support to further its awareness efforts and to expand its research program. Please contact us if you wish to help the Myocarditis Foundation save more lives from sudden cardiac death brought upon by the disease myocarditis.

The Myocarditis Foundation
3518 Echo Mountain Drive
Kingwood, Texas 77345
(281) 713-2962
www.myocarditisfoundation.org

The Myocarditis Foundation
Working to end sudden death from myocarditis through research, awareness, and advocacy
The Myocarditis Foundation is a 501(c)3 nonprofit organization dedicated to providing information about myocarditis to medical professionals, patients, and families with the goal of saving more lives.

Myocarditis is often misdiagnosed and is a poorly understood disease, which can rapidly progress to heart failure and death.

The Myocarditis Foundation works to fund research to find better ways to diagnose, treat and prevent myocarditis from having devastating effects on patients and their families around the world.

Since their establishment in 2005, the Myocarditis Foundation has awarded 15 research fellowship grants to young doctors in world-renowned university research facilities, with the shared goal of improving the care and medical outcome for patients affected by this devastating disease.

In addition to funding research, the Myocarditis Foundation takes great strides to increase the awareness of the disease myocarditis and its link to sudden cardiac related death in otherwise young, healthy people.

Who does the Myocarditis Foundation serve?

The Myocarditis Foundation provides information and support to patients diagnosed with myocarditis, families who have lost a loved one to myocarditis, and medical professionals who are seeking information for their patients with myocarditis.

How does the Myocarditis Foundation engage the medical community and the general public?

- Sponsors conferences, workshops, and lectures world-wide directed at patients, families, medical professionals, or professional heart disease groups
- Created a private and secure online research network to bring together and encourage the collaboration of researchers and physicians interested in and currently working on myocarditis research
- Maintains the only free and official myocarditis-specific website with features such as FAQ’s, research findings, blog, newsletters, and real-life stories

How does the Myocarditis Foundation spread awareness?

- Holds and supports fundraisers and awareness events around the world
- Sends myocarditis information to the medical community and the general public
- Uses newsletters, blogs, and social media to provide the most current information