The Myositis Association through funding from the Centers for Disease Control and Prevention is currently seeking adults and juveniles diagnosed with myositis to enroll in the Myositis Patient Registry, MYOVISION.

Registry surveys can be completed online or through a paper copy to be submitted by postal mail. All information collected will remain strictly confidential. The goal of MYOVISION is to provide a better understanding of potential triggers of the disease and characteristics of myositis patients. Analysis of the information collected and use of registry data will advance research and provide a better understanding of myositis.

**How Can You Participate in the Registry?**

- Contact The Myositis Association at 1-800-821-7356 or by email TMA@myositis.org to let us know of your interest. You can also go to the myositis home page www.myositis.org.

**Who Can Participate?**

- Anyone who was residing in the U. S. or Canada when diagnosed with myositis

**Why Should I Participate in the Registry?**

- To help physicians and researchers better understand this rare disease
- To identify environmental factors that may be related to the development of myositis
- To help encourage better treatments and further research toward a cure

For more information, contact The Myositis Association at:

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Email: tma@myositis.org
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