MS Minority Research

Accelerated Cure Project is interested in making sure that MS research benefits everyone. We have formed the MS Minority Research Engagement Partnership Network to understand how people with MS from different backgrounds and races think about medical research. The MS Minority Research Engagement Partnership Network is made up of people with multiple sclerosis, doctors, and other health leaders.

We want to understand how people with MS from different backgrounds and races think about participating in medical research. What do you think is important to study? What would help you be a part of the study?

If you are at least 21 years old and have multiple sclerosis, we invite you to complete a survey and share your thoughts on MS medical research. The survey should take around 15 minutes to complete. Everyone is invited!

Take the Survey

Conteste la Encuesta