The Muscular Dystrophy Association (MDA) is committed to transforming the lives of people with muscular dystrophy, ALS, and related neuromuscular diseases through innovations in science and care.

WHAT DO YOUR DOLLARS BUY?

$35 helps provide telehealth appointments at MDA Care Centers

$50 helps connect volunteers with MDA-funded clinical trials

$250 helps provide COVID-19 educational training for MDA providers

$500 helps fund vital research for a cure

WHAT IMPACT HAS COVID-19 HAD ON MDA?

MDA is an essential charity and we are at risk of losing over 30% of our funding due to the pandemic. We need your support so that MDA can continue to transform the lives for the 250,000 people living with neuromuscular disease. With your help, MDA can continue to fund groundbreaking research for promising treatments and provide families with the highest quality care from over 150+ MDA funded Care Centers in the United States.

HOW DOES MDA SPEND THE MONEY YOU RAISE?

72¢ helps support MDA’s efforts to:
- Fund groundbreaking research
- Educate and raise awareness
- Provide support and services to families across America

15¢ supports fundraising

13¢ supports overhead

WHAT PROGRAMS AND SERVICES DOES MDA OFFER TO FAMILIES?

- MDA hosts more than 3,000+ children with muscular dystrophy and related neuromuscular diseases at our nearly 54 week-long Summer Camps. This year, we invited more than 11,000 children to attend a Virtual Summer Camp program.
- 60,000+ children and adults receive comprehensive, expert care at 150 MDA Care Centers.
- MDA Advocates for access to care and screening for all neuromuscular diseases at the federal and state level.