

**We make every
day matter.**

**Because every day
matters to people
with Motor
Neurone Disease.**

About us

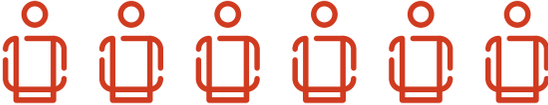
Every day we're here for people affected by Motor Neurone Disease (MND).

A diagnosis of MND brings home the preciousness of every day. So we do all we can to make every day count. We bring understanding and guidance. We deliver practical and financial support. We raise awareness and campaign for better care. We're not just here for now – as the UK's leading charity funder of MND research, we're striving for breakthroughs to develop new treatments and, ultimately, a cure.

MND moves fast. It takes away time, it takes away independence and it has no cure. Every piece of support, every research project, every pound raised, every kind word, and every day lived well, matters.

About Motor Neurone Disease

Today, six people



will be told they have MND and six people will die from the disease.

People with MND may lose their voice and movement.



And ultimately, their ability to breathe.

A third of people will die within a year of diagnosis.



More than half will die within two years.

MND is a fatal, rapidly progressing disease.



It affects the brain and spinal cord.

MND doesn't discriminate.



It affects people from all backgrounds and at all ages.

**There is no cure for MND.
Together we can change that. Your support matters.**
mndassociation.org

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Motor Neurone Disease Association

Francis Crick House

6 Summerhouse Road

Moulton Park

Northampton NN3 6BJ

Tel: 01604 250505

Email: enquiries@mndassociation.org

Website: mndassociation.org

Registered Charity no. 294354



Every day we support people affected by Motor Neurone Disease. Because with MND, every day matters.