Why join the registry?

- **To support** research projects by providing MTM and CNM data.
- **To be visible** for recruitment to research studies or clinical trials.
- **To encourage** further research into the conditions through the existence and availability of data.
- **To provide** information on living with MTM and CNM and develop standards of care.
- **To stay informed** about relevant news, research, or clinical trials.

We welcome registrations of:

- Individuals with a diagnosis of MTM or CNM (ideally confirmed via genetic testing or muscle biopsy).
- Female carriers of X-linked MTM.
- Deceased patients who had a confirmed diagnosis.

Anyone else who would like to receive newsletters or updates can request to join to our mailing list.

This international registry is available in 10 languages and is coordinated by the John Walton Muscular Dystrophy Research Centre at Newcastle University as part of the TREAT-NMD Network. Email or visit our website for more information.

🌐 [www.mtmcnmregistry.org](http://www.mtmcnmregistry.org) 📧 mtmcnmregistry@newcastle.ac.uk

**STRENGTH IN NUMBERS - REGISTER ONLINE TODAY**