

Importance of Timely and Accurate Diagnosis of Myotonic Disorders: Role of Electromyography

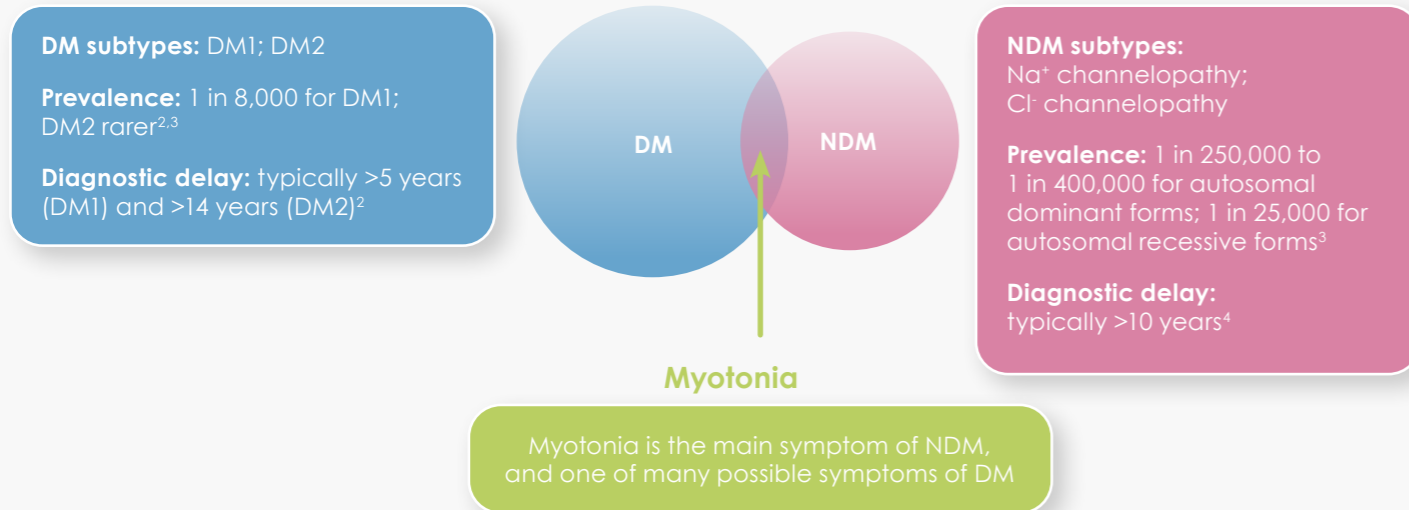
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Myotonic Dystrophy and Non-dystrophic Myotonias

- Myotonic disorders are a heterogeneous group of inherited neuromuscular disorders¹
- Myotonia is a symptom that is a common feature of several types and subtypes of myotonic disorders, including myotonic dystrophy (DM) and non-dystrophic (NDM) myotonias.
- Myotonia presents clinically as delayed muscle relaxation after voluntary contraction, leading to muscle stiffness or cramping, and/or electrophysiologically as spontaneous discharge of muscle fibres¹



Diagnostic delays

Variable, non-specific symptoms⁴⁻⁷

- Muscles affected
- Disease severity
- Age of onset
- Warm-up phenomenon
- Cold phenomenon
- Overlap with other diseases

Coping⁸

- A "family problem" (inherited disease) people manage by themselves

There are numerous reasons for delays in diagnosing DM and NDM

Lack of disease recognition⁴

- Rare diseases, not frequently encountered by non-specialist HCPs

Delays seeking medical help⁹

- Patients don't ask
- Non-specialists don't refer on

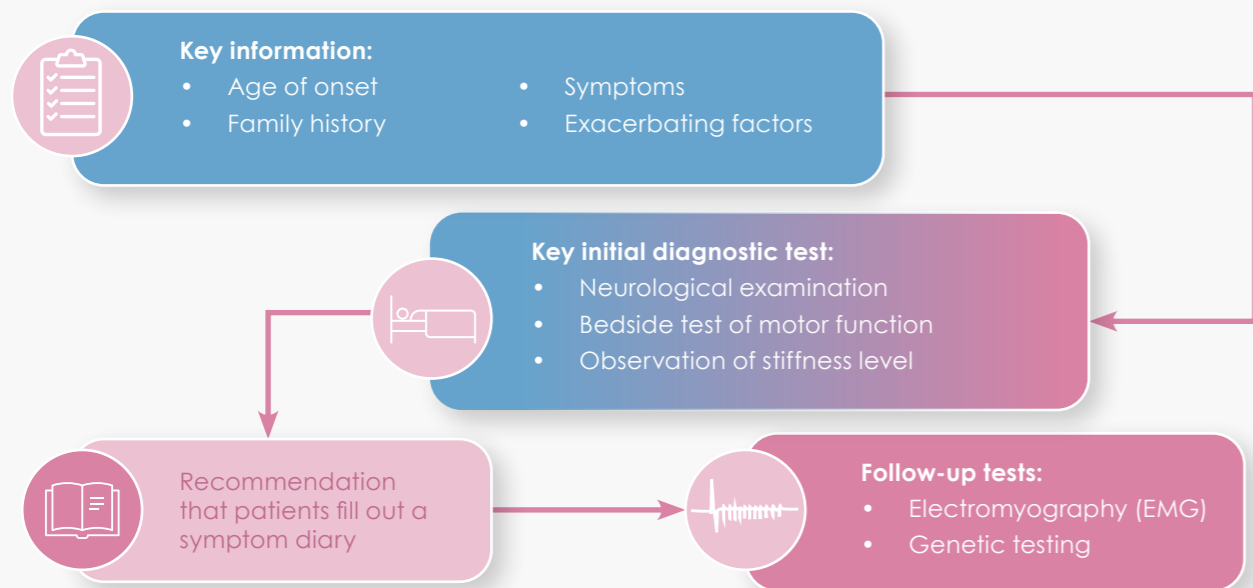
Under-recognised disease burden⁴

- Underestimation of impact on patients and carers

- Diagnostic delays have a negative impact on patients' wellbeing,² as they have to learn to cope with their condition, often by limiting what they do, instead of being offered treatment to ameliorate symptoms
- Timely and accurate diagnosis is important for genetic counselling and screening of systemic features in DM, as well as determining appropriate management¹
- Using EMG can help provide timely confirmation of a diagnosis

Diagnosis of Myotonic Disorders

Diagnostic Pathway (adapted from Stunnenberg et al 2020⁵)



Role of EMG in DM

Low need for EMG when there is a clear DM phenotype and clear signs of clinical myotonia

Greater need for EMG when:

- Genetic screening is required
- Myotonia fluctuates or is difficult to find

Role of EMG in NDM

- Useful when clinical myotonia is uncertain
- Helpful when interpreting relevance of a gene variant, and ensuring correct diagnosis
- Can prevent delay to appropriate medical management
- Can prevent erroneously offering inappropriate medical management

Abbreviations

DM: myotonic dystrophy; EMG: electromyography; HCP: healthcare professional; NDM: non-dystrophic myotonia.

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References:

1. Hahn C, Salajegheh MK. Myotonic disorders: a review article. Iran J Neurol. 2016;15(1):46-53.
2. Meola G. Clinical aspects, molecular pathomechanisms and management of myotonic dystrophies. Acta Myologica. 2013;32(3):154-65.
3. Lehmann-Horn F et al. Diagnostics and therapy of muscle channelopathies – guidelines of the Ulm Muscle Centre. Acta Myol. 2008;27(3):98-113.
4. Diaz-Manera J et al. Understanding the impact of non-dystrophic myotonia on patients and caregivers: results from a burden of disease healthcare survey. EMJ. 2021;6:2:37-46.
5. Stunnenberg BC et al. Guidelines on clinical presentation and management of nondystrophic myotonias. Muscle Nerve. 2020;62(4):430-44.
6. Matthews E et al. The non-dystrophic myotonias: molecular pathogenesis, diagnosis and treatment. Brain. 2010;133:9-22.
7. Hilbert JE et al. Diagnostic odyssey of patients with myotonic dystrophy. J Neurol. 2013;260(10):2497-504.
8. Trip J. Redefining the non-dystrophic myotonic syndromes: phenotypic characterisation based on genetic testing. Available at: <http://www.equipware.nl/mcbecker/myotonie%20van%20jeroen%20trip.pdf>. Last accessed: March 2024
9. Vereb N et al. Non-dystrophic myotonias: clinical and mutation spectrum of 70 German patients. J Neurol. 2021;268(5):1708-20.

Key Learnings

- EMG, while not necessary in every case, plays an important role in the timely diagnosis of both DM and NDM
- EMG, alongside genetic testing, can facilitate accurate differential diagnosis of disease subtypes