Rigid spine muscular dystrophy

Rigid spine muscular dystrophy (RSMD) is a form of congenital muscular dystrophy. This group of disorders cause muscle weakness and wasting (atrophy) beginning very early in life. In particular, RSMD involves weakness of the muscles of the torso and neck (axial muscles). Other characteristic features include spine stiffness and serious breathing problems.

In RSMD, muscle weakness is often apparent at birth or within the first few months of life. Affected infants can have poor head control and weak muscle tone (hypotonia), which may delay the development of motor skills such as crawling or walking. Over time, muscles surrounding the spine atrophy, and the joints of the spine develop deformities called contractures that restrict movement. The neck and back become stiff and rigid, and affected children have limited ability to move their heads up and down or side to side. Affected children eventually develop an abnormal curvature of the spine (scoliosis). In some people with RSMD, muscles in the inner thighs also atrophy, although it does not impair the ability to walk.

A characteristic feature of RSMD is breathing difficulty (respiratory insufficiency) due to restricted movement of the torso and weakness of the diaphragm, which is the muscle that separates the abdomen from the chest cavity. The breathing problems, which tend to occur only at night, can be life-threatening. Many affected individuals require a machine to help them breathe (mechanical ventilation) during sleep.

The combination of features characteristic of RSMD, particularly axial muscle weakness, spine rigidity, and respiratory insufficiency, is sometimes referred to as rigid spine syndrome. While these features occur on their own in RSMD, they can also occur along with additional signs and symptoms in other muscle disorders. The features of rigid spine syndrome typically appear at a younger age in people with RSMD than in those with other muscle disorders.

Frequency

RSMD is thought to be a rare disorder, although its prevalence is unknown.

Causes

Mutations in a gene called \( \text{SELENON} \) (formerly \( \text{SEPN1} \)) cause about 40 percent of cases of RSMD. When caused by mutations in this gene, the condition is known as rigid spine muscular dystrophy 1 (RSMD1). The genetic cause of the remainder of cases is unknown.

The \( \text{SELENON} \) gene provides instructions for making a protein known as selenoprotein N. The specific job of selenoprotein N is unknown, but researchers suspect it plays a
role in the formation of muscle tissue before birth. It may also be important for normal muscle function after birth. The gene mutations that cause RSMD1 are thought to reduce the amount of selenoprotein N or impair its activity in cells. It is unclear how a shortage of working selenoprotein N leads to muscle weakness and other features of RSMD1.

RSMD1 is part of a spectrum of muscle disorders called SELENON-related (or SEPN1-related) myopathy, which also includes the classic form of multiminicore disease, desmin-related myopathy with Mallory body-like inclusions, and a small subset of cases of congenital fiber-type disproportion. While these other disorders share the characteristic features of RSMD1, they each also involve distinctive abnormalities of the muscle fibers that can only be seen when viewed under a microscope. Because these conditions have a similar pattern of signs and symptoms and are caused by mutations in the same gene, many researchers believe that they are all part of a single syndrome with variable signs and symptoms. It is unclear why mutations in the SELENON gene cause the different muscle fiber abnormalities that distinguish the separate conditions.

Inheritance Pattern

This condition is inherited in an autosomal recessive pattern, which means both copies of the gene in each cell have mutations. The parents of an individual with an autosomal recessive condition each carry one copy of the mutated gene, but they typically do not show signs and symptoms of the condition.

Other Names for This Condition

- congenital muscular dystrophy with spine rigidity syndrome
- muscular dystrophy, congenital, merosin-positive, with early spine rigidity
- rigid spinal muscular dystrophy
- rigid spine congenital muscular dystrophy
- RSMD

Diagnosis & Management

Genetic Testing Information

- What is genetic testing? /primer/testing/genetictesting
- Genetic Testing Registry: Eichsfeld type congenital muscular dystrophy

Research Studies from ClinicalTrials.gov

- ClinicalTrials.gov
  https://clinicaltrials.gov/ct2/results?cond=%22rigid+spine+muscular+dystrophy+1%22+OR+%22rigid+spine+muscular+dystrophy%22
Additional Information & Resources

Health Information from MedlinePlus

- Encyclopedia: Scoliosis
  https://medlineplus.gov/ency/article/001241.htm
- Health Topic: Muscular Dystrophy
  https://medlineplus.gov/musculardystrophy.html

Genetic and Rare Diseases Information Center

- Rigid spine syndrome
  https://rarediseases.info.nih.gov/diseases/4723/rigid-spine-syndrome

Additional NIH Resources

- National Institute of Neurological Disorders and Stroke: Muscular Dystrophy: Hope Through Research

Educational Resources

- MalaCards: rigid spine muscular dystrophy 1
  https://www.malacards.org/card/rigid_spine_muscular_dystrophy_1
- Neuromuscular Disease Center, Washington University
  https://neuromuscular.wustl.edu/syncm.html#cmspine
- Orphanet: Rigid spine syndrome
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=97244

Patient Support and Advocacy Resources

- Cure CMD: SEPN1-related myopathy
  https://www.curecmd.org/sepn1
- Muscular Dystrophy Association: Congenital Muscular Dystrophy
  https://www.mda.org/disease/congenital-muscular-dystrophy
- Muscular Dystrophy UK
  https://www.musculardystrophyuk.org/about-muscle-wasting-conditions/congenital-muscular-dystrophy-cmd/sepn1-related-myopathy-factsheet/
- Resource List from the University of Kansas Medical Center: Muscular Dystrophy / Atrophy
  http://www.kumc.edu/gec/support/muscular.html
Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28rigid+spine+muscular+dystrophy%5BTIAB%5D%29+OR+%28SEPN1-related+muscular+dystrophy%5BTIAB%5D%29+OR+%28SELENON-related+muscular+dystrophy%5BTIAB%5D%29+OR+%28congenital+muscular+dystrophy+with+rigid+spine%5BTIAB%5D%29+OR+%28rigid+spine+muscular+dystrophy+1%5BTIAB%5D%29%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+3600+days%22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

- RIGID SPINE MUSCULAR DYSTROPHY 1
  http://omim.org/entry/602771

Medical Genetics Database from MedGen

- Rigid spine syndrome

Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/26780752

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/28606403

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21131290

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/22527882

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/23481446

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11528383
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/10545040

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21670436

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