

38,400 Results Any time ▾

[A missense mutation in DYNC1H1 gene causing spinal ...](#)<https://www.sciencedirect.com/science/article/pii/S0028384317303468>

Mar 01, 2018 · SMALED caused by DYNC1H1 mutations is not a fatal condition and the lack of autopsy examination makes it difficult to prove these hypotheses. There are other phenotypes that have been...

Cited by: 3

Author: Joyutpal Das, James B. Lilleker, Kavaldeep ...

Publish Year: 2017

[DYNC1H1 - an overview | ScienceDirect Topics](#)<https://www.sciencedirect.com/.../dync1h1>

Shortly after the initial description of dynein mutations with a phenotype of intellectual disability, an analysis of a large pedigree with dominant axonal CMT disease, type 2 determined that the causative...

PEOPLE ALSO ASK

What are the effects of mutations in DYNC1H1?



Is there a heterozygous mutation in dynein heavy chain 1?



How are mutations in NDE1 related to microcephaly?



Are there any recessive mutations in the NDE1 gene?



Feedback

[Myasthenic syndrome, congenital, 18 \(Concept Id: C4225364\)](#)<https://www.ncbi.nlm.nih.gov/medgen/906793>

From OMIM Congenital myasthenic syndrome 18 is an autosomal dominant presynaptic neuromuscular disorder characterized by early-onset muscle weakness and easy fatigability associated with delayed...

[Arts syndrome with a novel missense mutation in the PRPS1 ...](#)<https://pubmed.ncbi.nlm.nih.gov/27256512>

Arts syndrome is characterized by early-onset hypotonia, ataxia, intellectual disability, sensorineural hearing impairment, progressive optic atrophy, and a tendency to develop infections. Arts syndrome is a...

Cited by: 0

Author: Kunihi Maruyama, Shunsuke Onawa, Naoko

[A missense mutation in DYNC1H1 gene causing spinal ...](#)

<https://www.sciencedirect.com/science/article/pii/S0028384317303468>

Mar 01, 2018 · SMALED caused by DYNC1H1 mutations is not a fatal condition and the lack of autopsy examination makes it difficult to prove these hypotheses. There are other phenotypes that have been associated with DYNC1H1 mutations. Weedon et al. reported DYNC1H1 mutations in a large family with Charcot Marie Tooth type 2 . Early onset, **delayed motor development**, distal lower limb wasting and **weakness** ...

Cited by: 3 Author: Joyutpal Das, James B. Lilleker, Kavaldeep ...

Publish Year: 2017

[\(PDF\) A Novel De Novo Variant in DYNC1H1 Causes Spinal ...](#)

<https://www.researchgate.net/publication/353408025...>

Jul 01, 2021 · This **report** provides important clinical evidence indicating the de novo heterozygous missense mutation c.751C>T in the DYNC1H1 gene is pathogenic causing SMALED. **Muscle** ...

Search Tools

Turn off Hover Translation (关闭取词)


People also ask

- What are the effects of mutations in DYNC1H1? ▾
- Which is heterozygous mutation in dynein heavy chain 1? ▾
- How are mutations in NDE1 related to microcephaly? ▾

30-Jul-2021 11:11AM

2541 words • 5 matches • 4 spaces

FAQ

 iThenticate®

68893-crosscheck.docx

Quotient Excluded
Bibliography Excluded
3%
(26/10)

Match Overview

1

Crossref 35 words
Salmon, A. "Rapid Development of Post-radiotherapy Sarcoma and Breast Cancer in a Patient with a Novel Gemin3 Deletion"

1%

2

Crossref 15 words
Jeyaraj, S., Sotgiu, M., Mages, M., Enik, E., P. Hoffman, Sebastian, C., Carolina Test-Rocha, "Exome Sequencing"

1%

3

Crossref 12 words
Sami Rizzari, Abdul Razak Haniff, Prithvi Nar, Madhva Mohamed, Fatma Saeed, "Characterization of an Enzyme"

<1%

4

Internet 12 words
crawled on 23-Mar-2021
path-csa.them.ac.uk

<1%

A missense mutation in *DYNC1H1* gene caused psychomotor developmental delay and muscle weakness: A case report

Ding FJ *et al.* A missense mutation in *DYNC1H1* gene

Feng-Juan Ding, Cui-Zhen Lyu, Wei Zhang, Hua Jin

Received: June 7, 2021

Revised: July 9, 2021

Accepted:

Published online:

PDF, 1 OF 10

Test-Only Report

国内版

国际版

Missense mutation in DYNC1H1 gene caused psychomotor develo



ALL

IMAGES

VIDEOS

9,150 Results

Any time ▾

Including results for missense mutation in dync1h1 gene **causes** psychomotor developmental delay and muscle weakness a case report.

Do you want results only for Missense mutation in DYNC1H1 gene caused psychomotor developmental delay and muscle weakness: A case report?

A missense mutation in DYNC1H1 gene causing spinal ...

<https://www.sciencedirect.com/science/article/pii/S0028384317303468>

Mar 01, 2018 · Here we report a family with SMALED caused by a pathogenic heterozygous missense c.1809 A > T, p.glu603Asp mutation in DYNC1H1. The main clinical features were congenital hip displacement, talipes, delayed motor development, wasting and weakness in lower limbs with relative sparing of upper extremities and very slow disease progression.

Cited by: 3

Author: Joyutpal Das, James B. Lilleker, Kavalde...

Publish Year: 2017

(PDF) A Novel De Novo Variant in DYNC1H1 Causes Spinal ...

<https://www.researchgate.net/publication/353408025...>

This report provides important clinical evidence indicating the de novo heterozygous missense mutation c.751C>T in the DYNC1H1 gene is pathogenic causing SMALED. Muscle MRI is more specific than ...

PEOPLE ALSO ASK

What are the effects of mutations in DYNC1H1?



Which is heterozygous mutation in dynein heavy chain 1?



How are mutations in NDE1 related to microcephaly?



How many chains are there in the DYNC1H1 homodimer?



Feedback

Mutations in DYNC1H1 cause severe intellectual disability ...

<https://www.researchgate.net/publication/221864565...>

Autosomal dominant mutations of DYNC1H1 cause a range of neurogenetic diseases, including: