



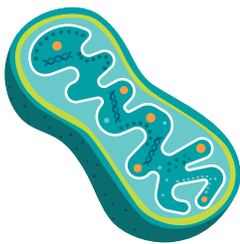
# Understanding

# TK2d

## A Rare, Under-Recognized Genetic Mitochondrial Disorder

### THYMIDINE KINASE 2 DEFICIENCY

Thymidine kinase 2 deficiency (TK2d) is a debilitating and life-threatening genetic disorder that causes progressive and severe muscle weakness.<sup>1,2,3,4</sup> Many patients lose the ability to walk, eat, and breathe independently.<sup>1,2,3,4</sup>



#### What causes TK2d?

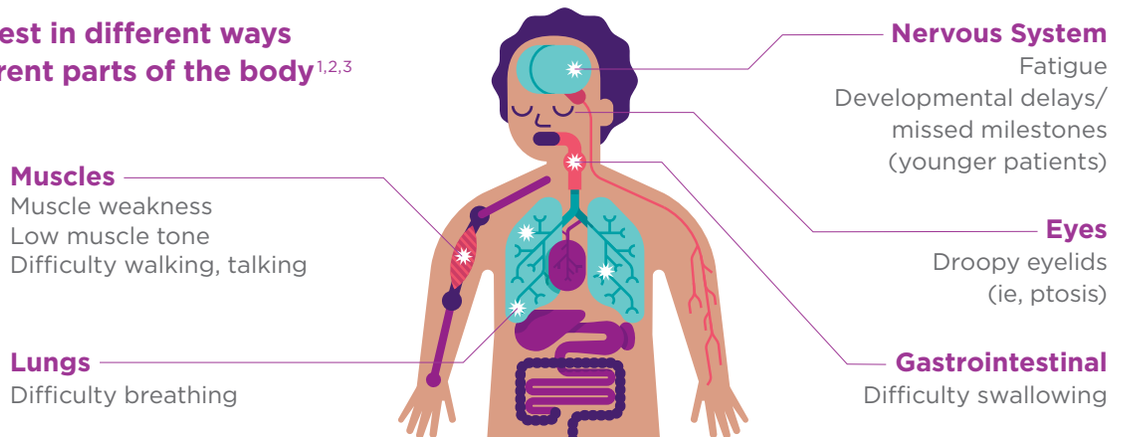
TK2d is caused by a genetic mutation in the TK2 gene.<sup>1,4</sup> This mutation leads to a decrease in mitochondrial DNA production.<sup>1,4</sup> Mitochondria generate most of the energy that powers our cells. Errors in mitochondrial DNA can lead to insufficient energy, and as a result, muscles and organs can't function properly.<sup>4,5,6</sup>

#### How many people have TK2d?

1 in 5,000 people have some form of genetic mitochondrial disease.<sup>7</sup> Prevalence of TK2d itself is still being researched.

## PROGRESSIVELY WORSENING SYMPTOMS

TK2d can manifest in different ways and affect different parts of the body<sup>1,2,3</sup>



## IMPACTS ALL AGES: Symptoms can progress slowly or rapidly<sup>1,2</sup>

EARLY ONSET: OFTEN MORE SEVERE

LATER ONSET: OFTEN LESS SEVERE



**Normal Development**



**Infant/Child Onset**



**Adolescent Onset**



**Adult Onset**

## TK2D MANAGEMENT TODAY: No FDA-approved therapies currently available<sup>8</sup>

### Patients require medical equipment and therapies<sup>1,3,8</sup>

- Respiratory support
- Feeding tubes
- Wheelchairs
- Respiratory physiotherapy
- Physical therapy

Patients may take nutraceuticals (“mito-cocktail”) to manage disease symptoms<sup>8</sup>



### Patients rely on supportive care provided by multi-disciplinary team<sup>7</sup>

- Neurologists
- Pulmonologists
- Metabolic specialists
- Gastroenterologists
- Physical therapists
- Speech therapists
- Clinical geneticists
- Genetic counselors



## Genetic Testing Is Necessary

**Earlier accurate diagnosis helps to identify patients sooner and get them on supportive care and into clinical trials faster.**<sup>1,2,3</sup>

Because TK2d can present like other diseases (e.g. muscular dystrophy, Pompe, SMA, mtDNA depletion syndrome, and others), genetic testing is needed to confirm a diagnosis.<sup>1</sup>

**If you or your family member has TK2d:**

Learn more at [www.tk2d.com](http://www.tk2d.com)

## Our Commitment

At Zogenix, we are proud to partner with physicians and patient communities in our work.

**Together we can bring hope and support to patients and families impacted by rare diseases.**



**ZOGENIX**  
Rare Disease Therapies

<sup>1</sup> Garone C, Taylor RW, Nascimento A, et al. Retrospective natural history of thymidine kinase 2 deficiency. *J Med Genet.* 2018;55(8):515-21. <sup>2</sup> Wang J, Kim E, Dai H, et al. Clinical and molecular spectrum of thymidine kinase 2-related mtDNA maintenance defect. *Mol Genet Metab.* 2018;124:124-30. <sup>3</sup> Domínguez-González C, Hernández-Lain A, Rivas E, et al. Late-onset thymidine kinase 2 deficiency: a review of 18 cases. *Orphanet J Rare Dis.* 2019;14(1):100. <sup>4</sup> National Institute of Health. TK2-related mitochondrial DNA depletion syndrome, myopathic form. <https://medlineplus.gov/genetics/condition/tk2-related-mitochondrial-dna-depletion-syndrome-myopathic-form/#genes>. Accessed April 27, 2021. <sup>5</sup> United Mitochondrial Disease Foundation. Understanding & Navigating Mitochondrial Disease. <https://www.umdf.org/what-is-mitochondrial-disease-2/>. Accessed April 27, 2021. <sup>6</sup> Hirano M, Marti R, Ferreiro-Barros C, et al. Defects of intergenomic communication: autosomal disorders that cause multiple deletions and depletion of mitochondrial DNA. *Semin Cell Dev Biol.* 2001;12:417-27. <sup>7</sup> Parikh S, Goldstein A, Karaa A, et al. Patient care standards for primary mitochondrial disease: a consensus statement from the Mitochondrial Medicine Society. *Genet Med.* 2017;19(12):1010-38. <sup>8</sup> El-Hattab AW and Scaglia F. Mitochondrial DNA depletion syndromes: review and updates of genetic basis, manifestations, and therapeutic options. *Neurotherapeutics.* 2013;10:186-98.