

# **Novel compound heterozygous mutations in TTI2 cause syndromic intellectual disability in a Chinese family**

Wang R, Han S, Liu H, Khan A, Xiaerbati H, Yu X, Huang J, Zhang X.

Frontiers in genetics

2019; 10:e1060

## **ARTICLE IDENTIFIERS**

DOI: 10.3389/fgene.2019.01060

PMID: 31737043

PMCID: PMC6830114

## **JOURNAL IDENTIFIERS**

LCCN: not available

pISSN: not available

eISSN: 1664-8021

OCLC ID: 731654963

CONS ID: not available

US National Library of Medicine ID: 101560621

This article was identified from a query of the SafetyLit database.