

# Resource Summary Report

Generated by [dkNET](#) on Sep 19, 2026

## GM04026

RRID:CVCL\_AX77

Type: Cell Line

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### Proper Citation

RRID:CVCL\_AX77

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### Cell Line Information

**URL:** [https://web.expasy.org/cellosaurus/CVCL\\_AX77](https://web.expasy.org/cellosaurus/CVCL_AX77)

**Proper Citation:** RRID:CVCL\_AX77

**Description:** Cell line GM04026 is a Finite cell line with a species of origin Homo sapiens (Human)

**Sex:** Male

**Disease:** Fragile X syndrome

**Comments:** Population: Caucasian.

**Category:** Finite cell line

**Organism:** Homo sapiens (Human)

**Name:** GM04026

**Cross References:** CLO:CLO\_0016230, Coriell:GM04026, Wikidata:Q54838375

**ID:** CVCL\_AX77

**Record Creation Time:** 20220427T215933+0000

**Record Last Update:** 20260913T003236+0000

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### Ratings and Alerts

No rating or validation information has been found for GM04026.

No alerts have been found for GM04026.

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### Data and Source Information

**Source:** [Cellosaurus](#)

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## Usage and Citation Metrics

We found 1 mentions in open access literature.

**Listed below are recent publications.** The full list is available at [dkNET](#) .

Susco SG, et al. (2022) Molecular convergence between Down syndrome and fragile X syndrome identified using human pluripotent stem cell models. Cell reports, 40(10), 111312.