

Resource Summary Report

Generated by [RRID](#) on Sep 3, 2026

[FINi002-A](#)

RRID:CVCL_C8H8

Type: Cell Line

Proper Citation

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Cell Line Information

URL: https://web.expasy.org/cellosaurus/CVCL_C8H8

Proper Citation: RRID:CVCL_C8H8

Description: Cell line FINi002-A is a Induced pluripotent stem cell with a species of origin Homo sapiens (Human)

Sex: Male

Disease: Parkinson disease 2, autosomal recessive juvenile

Defining Citation: [PMID:37890334](#)

Comments: Population: Caucasian., From: The Florey Institute of Neuroscience and Mental Health, University of Melbourne; Melbourne; Australia.

Category: Induced pluripotent stem cell

Organism: Homo sapiens (Human)

Name: FINi002-A

Synonyms: FI.CPLT.PRKN.R275W+del_e8.PK006

Cross References: BioSamples:SAMEA112835223, hPSCreg:FINi002-A, Wikidata:Q123031280

ID: CVCL_C8H8

Record Creation Time: 20231016T222434+0000

Record Last Update: 20260830T000720+0000

Ratings and Alerts

No rating or validation information has been found for FINi002-A.

No alerts have been found for FINi002-A.

Data and Source Information

Source: [Cellosaurus](#)

Usage and Citation Metrics

We found 1 mentions in open access literature.

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

Pavan C, et al. (2023) Generation of the iPSC line FINi002-A from a male Parkinson's disease patient carrying compound heterozygous mutations in the PRKN gene. Stem cell research, 73, 103211.