

# Resource Summary Report

Generated by [RRID](#) on Aug 28, 2026

## CPGHi002-A

RRID:CVCL\_A1AZ

Type: Cell Line

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### Proper Citation

RRID:CVCL\_A1AZ

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### Cell Line Information

**URL:** [https://web.expasy.org/cellosaurus/CVCL\\_A1AZ](https://web.expasy.org/cellosaurus/CVCL_A1AZ)

**Proper Citation:** RRID:CVCL\_A1AZ

**Description:** Cell line CPGHi002-A is a Induced pluripotent stem cell with a species of origin Homo sapiens (Human)

**Sex:** Female

**Disease:** Autosomal dominant congenital deafness with onychodystrophy

**Defining Citation:** [PMID:32961450](#), [PMID:33714068](#)

**Comments:** Population: Chinese; Han., From: Chinese PLA Institute of Otolaryngology, Chinese PLA General Hospital; Beijing; China.

**Category:** Induced pluripotent stem cell

**Organism:** Homo sapiens (Human)

**Name:** CPGHi002-A

**Synonyms:** JYIPS0056

**Cross References:** BioSamples:SAMEA7340582, hPSCreg:CPGHi002-A, Wikidata:Q102113716

**ID:** CVCL\_A1AZ

**Record Creation Time:** 20220427T215517+0000

**Record Last Update:** 20260815T232120+0000

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### Ratings and Alerts

No rating or validation information has been found for CPGHi002-A.

No alerts have been found for CPGHi002-A.

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## Data and Source Information

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

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## Usage and Citation Metrics

We have not found any literature mentions for this resource.