

# Resource Summary Report

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## The NINDS Human Cell and Data Repository (NHCDR)

RRID:SCR\_016319

Type: Tool

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### Proper Citation

The NINDS Human Cell and Data Repository (NHCDR) (RRID:SCR\_016319)

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### Resource Information

**URL:** <https://stemcells.nindsgenetics.org/>

**Proper Citation:** The NINDS Human Cell and Data Repository (NHCDR) (RRID:SCR\_016319)

**Description:** Cell sources currently include fibroblasts and/or induced pluripotent stem cells for Alzheimer's Disease, Amyotrophic Lateral Sclerosis (ALS), Ataxia-telangiectasia, Frontotemporal Lobar Degeneration (FTD), Huntington's Disease, Parkinson's Disease, and healthy controls. Cell sources, including isogenic cell lines for current and new diseases covered by the NINDS will be added over the next several years.

**Abbreviations:** NHCDR

**Synonyms:** NINDS Human Cell and Data Repository (NHCDR)

**Resource Type:** tissue bank, biomaterial supply resource, material resource

**Keywords:** Stem, cell, fibroblast, pluripotent, isogenic

**Related Condition:** Alzheimer's Disease, Amyotrophic Lateral Sclerosis (ALS), Ataxia-telangiectasia, Frontotemporal Lobar Degeneration (FTD), Huntington's Disease, Parkinson's Disease

**Funding:** NLM ;  
NINDS

**Availability:** Restricted

**Resource Name:** The NINDS Human Cell and Data Repository (NHCDR)

**Resource ID:** SCR\_016319

**Alternate URLs:** <https://nindsgenetics.org/>

**Record Creation Time:** 20220129T080330+0000

**Record Last Update:** 20260804T043622+0000

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## Ratings and Alerts

No rating or validation information has been found for The NINDS Human Cell and Data Repository (NHCDR).

No alerts have been found for The NINDS Human Cell and Data Repository (NHCDR).

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## Data and Source Information

**Source:** [SciCrunch Registry](#)

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## Usage and Citation Metrics

We found 16 mentions in open access literature.

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

Drwesh L, et al. (2025) Methodological validation of Miro1 retention as a candidate Parkinson's disease biomarker. NPJ Parkinson's disease, 11(1), 270.

Baninameh Z, et al. (2025) Development and validation of a sensitive sandwich ELISA against human PINK1. Autophagy, 21(5), 1144.

Jiang Q, et al. (2025) Robust differentiation of NK cells from MSLN.CAR-IL-15-engineered human iPSCs with enhanced antitumor efficacy against solid tumors. Science advances, 11(18), eadt9932.

Kim KH, et al. (2024) Posttranscriptional regulation of FAN1 by miR-124-3p at rs3512 underlies onset-delaying genetic modification in Huntington's disease. Proceedings of the National Academy of Sciences of the United States of America, 121(16), e2322924121.

DiFiglia M, et al. (2024) Towards Standardizing Nomenclature in Huntington's Disease Research. Journal of Huntington's disease, 13(2), 119.

Yu SF, et al. (2024) Protein Assembly Modulation: A New Approach to Amyotrophic Lateral Sclerosis (ALS) Therapeutics. Journal of experimental neurology, 5(4), 210.

Cecerska-Hery? E, et al. (2023) The Use of Stem Cells as a Potential Treatment Method for Selected Neurodegenerative Diseases: Review. Cellular and molecular neurobiology, 43(6), 2643.

Bharat V, et al. (2023) A mitochondrial inside-out iron-calcium signal reveals drug targets for Parkinson's disease. Cell reports, 42(12), 113544.

Tamiz AP, et al. (2022) A focus on the neural exposome. *Neuron*, 110(8), 1286.

Kühn R, et al. (2021) Human Induced Pluripotent Stem Cell Models of Frontotemporal Dementia With Tau Pathology. *Frontiers in cell and developmental biology*, 9, 766773.

, et al. (2021) An integrated multi-omic analysis of iPSC-derived motor neurons from C9ORF72 ALS patients. *iScience*, 24(11), 103221.

Patel A, et al. (2020) Establishment and characterization of two iPSC lines derived from healthy controls. *Stem cell research*, 47, 101926.

Schwartzentruber A, et al. (2020) Oxidative switch drives mitophagy defects in dopaminergic parkin mutant patient neurons. *Scientific reports*, 10(1), 15485.

Ng SS, et al. (2018) Human iPS derived progenitors bioengineered into liver organoids using an inverted colloidal crystal poly (ethylene glycol) scaffold. *Biomaterials*, 182, 299.

Wiatr K, et al. (2018) Huntington Disease as a Neurodevelopmental Disorder and Early Signs of the Disease in Stem Cells. *Molecular neurobiology*, 55(4), 3351.

Dunås T, et al. (2017) A Stereotactic Probabilistic Atlas for the Major Cerebral Arteries. *Neuroinformatics*, 15(1), 101.