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Lab resource: Stem Cell Line

Generation of a human iPSC line, INMi002-A, carrying the most prevalent *USH2A* variant associated with Usher syndrome type 2



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ABSTRACT

We generated an induced pluripotent stem cell (iPSC) line using dermal fibroblasts from a patient with Usher syndrome type 2 (USH2). This individual was homozygous for the most prevalent variant reported in the *USH2A* gene, c.2299delG localized in exon 13. Reprogramming was performed using the non-integrative Sendai virus reprogramming method and the human OSKM transcription factor cocktail under feeder-free culture conditions. This iPSC line will be an invaluable tool for studying the pathophysiology of USH2 and for testing the efficacy of novel treatments.

Resource table

Unique stem cell line identifier	INMi002-A		
Alternative name(s) of stem cell line	USH2A-USH-iPSC		
Institution	Institute for Neurosciences of Montpellier,		
	Montpellier, France		
Contact information of distributor	Vasiliki Kalatzis vasiliki.kalatzis@inserm.		
	fr		
Type of cell line	iPSC		
Origin	Human		
Additional origin info	Age: 59 years old		
Ū.	Sex: Female		
	Ethnicity: Caucasian		
Cell Source	Dermal fibroblasts		
Clonality	Clonal		
Method of reprogramming	Non-integrative Sendai virus vectors		
Genetic modification	Yes		
Type of modification	Congenital mutation		
Associated disease	Usher Syndrome type II		
Gene/locus	USH2A, 1q41		
Method of modification	N/A		
Name of transgene or resistance	N/A		
Inducible/constitutive system	N/A		
Date archived/stock date	December 2017		
Cell line repository/bank	https://hpscreg.eu/cell-line/INMi002-A		
Ethical approval	Regional committee: CPP Southern		
	Mediterranean I (2014-A00549-38)		
	National committee: ANSM (140549B-62)		

Resource utility

We established an iPSC line from a patient with Usher syndrome type 2 (USH2), characterized by retinitis pigmentosa and hearing loss, homozygous for the recurrent *USH2A* variant, c.2299delG. This line will allow modelling of the USH2 retinal and inner ear defects, and the development of novel gene and cell therapies.

Resource details

Mutations in *USH2A* cause a syndromic inherited retinal dystrophy (IRD) known as Usher syndrome type 2 (USH2), which is characterized by progressive hearing and vision loss. In addition, mutations in *USH2A* also cause a wide majority of autosomal recessive retinitis pigmentosa (RP) cases (Kremer et al., 2006). In the present study, we have generated an iPSC line from a patient with USH2 carrying a homozygous variant in exon 13 of the *USH2A* gene, c.2299delG; p.Glu767Serfs*21. This is a recurrent mutation originating from a common ancestor. Therefore, c.2299delG is observed more frequently in the population compared to the > 600 *USH2A* mutations identified (Lenassi et al., 2015), conferring a great deal of interest from a clinical perspective.

Human dermal fibroblasts were isolated and cultured from a patient skin biopsy sample and reprogrammed using an integration-free method, the CytoTune[™]-iPS 2.0 Sendai Reprogramming Kit. This method is based on transient overexpression of the four Yamanaka factors: *OCT3/4*, *SOX2*, *KLF4* and *c-MYC* (Takahashi et al., 2007). The morphology of the iPSC line generated displayed a typical colony

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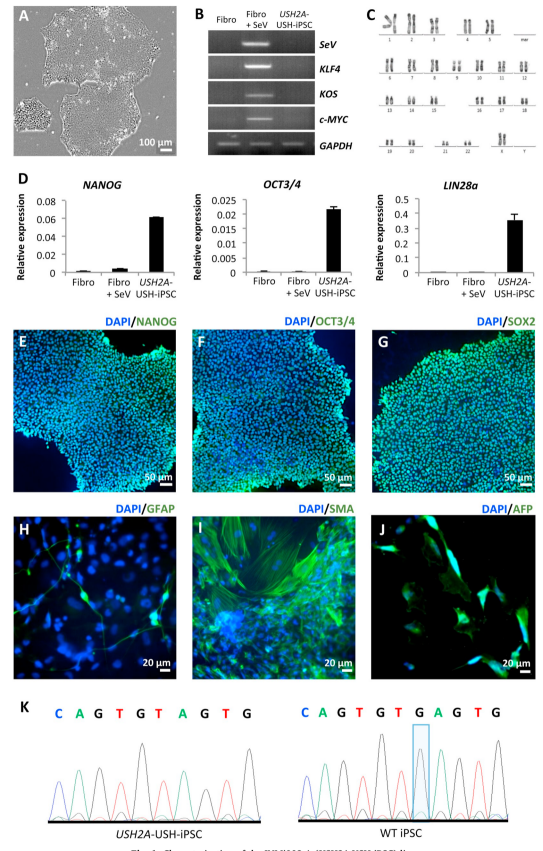


Fig. 1. Characterization of the INMi002-A (USH2A-USH-iPSC) line.

Table 1

Characterization and validation.

Classification	Test	Result	Data	
Morphology	Photography	Normal morphology	Fig. 1A	
	Microscopy			
Phenotype	Qualitative analysis	Positive for pluripotency markers: NANOG, OCT3/4 and SOX2	Fig. 1E–1G	
	Immunocytochemistry			
	Quantitative analysis qPCR	Expression of pluripotency markers: NANOG, OCT3/4, LIN28a	Fig. 1D	
Genotype	Karyotype (G-banding) and resolution	46XX Resolution 400-500	Fig. 1C	
Identity	Microsatellite PCR (mPCR) OR	DNA Profiling Performed	Available with authors	
		D3S306		
		D8S532		
		D11S4191		
		D17S787		
		D19S572		
	STR analysis	Not performed	N/A	
Mutation analysis	Sequencing	Homozygous, USH2A c.2299delG	Fig. 1K	
	Southern Blot OR WGS	Not performed	N/A	
Microbiology and virology	Mycoplasma	Mycoplasma testing by luminescence. Negative	Supplementary File 1	
Differentiation potential	Embryoid body formation	Expression of GFAP (ectoderm), SMA (mesoderm) and AFP (endoderm) in iPSC-derived EBs	Fig. 1H–1J	
Donor screening (OPTIONAL)	Hepatitis A, Hepatitis B, Hepatitis C, HIV 1–2	Negative	Available with authors	
Genotype additional info	Blood group genotyping	N/A	N/A	
(OPTIONAL)	HLA tissue typing	N/A	N/A	

appearance comprised of tightly packed cells (Fig. 1A). The absence of the Sendai reprogramming vectors in the INMi002-A clones was confirmed using reverse transcription (RT)-PCR. As a negative control, nontransduced patient fibroblasts (Fibro) did not carry the Sendai vectors. By contrast, the transduced fibroblasts (Fibro + SeV) expressed all three vectors that were used for reprogramming. Due to their non-integrative nature, the INMi002-A cell line had lost these vectors by passage 12 (P12) (Fig. 1B). The iPSC line generated showed a normal 46,XX karyotype at P12, which excluded major chromosomal abnormalities as a result of the reprogramming process (Fig. 1C).

Using real time polymerase chain reaction (qPCR), we showed expression of the endogenous pluripotency markers, NANOG, OCT3/4 and LIN28a in the INMi002-A cell line when compared to non-transduced (Fibro) or transduced (Fibro + SeV; Fig. 1D) fibroblasts. The pluripotency state was further confirmed by immunofluorescence staining of NANOG (Fig. 1E), OCT3/4 (Fig. 1F) and SOX2 (Fig. 1G). The ability of the INMi002-A cell line to give rise to the three embryonic cell layers was determined by spontaneous differentiation of the iPSC into embryoid bodies (EBs). EBs were plated onto Matrigel-coated dishes and the expression of the three germ layer markers, Glial Fibrillary Acidic Protein (GFAP; ectoderm; Fig. 1H), Smooth Muscle Actin (SMA; mesoderm; Fig. 1I) and α-Fetoprotein (AFP; endoderm; Fig. 1J) was verified by immunofluorescence staining. We verified the presence of the homozygous causative mutation in exon 13 of USH2A (c.2299delG) in the INMi002-A iPSC, as compared to wild type iPSC, by Sanger sequencing (Fig. 1K). The identity of the patient iPSC line was confirmed by microsatellite PCR analysis in comparison to fibroblasts of the same individual and wild type iPSC (available with the authors). Lastly, the generated INMi002-A cell line was confirmed to be free of mycoplasma contamination (Supplementary File 1).

Materials and methods

Human dermal fibroblast cell culture

Dermal fibroblasts derived from a skin biopsy were cultured in AmnioMAX C100 basal media with GlutaMAX (Gibco) containing 10% decomplemented foetal calf serum (FCS; Lonza), 1% penicillin-streptomycin-amphotericin B (Lonza) and 2% AmnioMax-C100 supplement (Gibco).

iPSC generation

Fibroblasts were seeded and reprogrammed using the integrationfree CytoTuneTM-iPS 2.0 Sendai Reprogramming Kit as previously described (Torriano et al., 2017). iPSC colonies were cultured in Essential 8 (E8) medium (Gibco) under 5% CO₂ at 37 °C, and passaged (1:10 dilution; twice weekly) using Versene (Gibco).

Karyotype analysis

Preparation of the iPSC for karyotype analysis was performed as previously described without modification (Torriano et al., 2017). Twenty metaphase spreads were counted and analyses were performed using standard G-banding procedures by the Chromostem facility (CHU Montpellier, France).

RT-PCR and qPCR analysis

RNA was extracted using the RNeasy Mini Kit (QIAGEN) and cDNA synthesis performed using the SuperScript[®] III First-Strand Synthesis System (Life Technologies) and random hexamers (Life Technologies), according to the manufacturer's protocols. RT-PCR was performed using a standard protocol on an Applied Biosystems Veriti 96-well thermal cycler. qPCR analysis was performed using the FastStart SYBR Green I Master mix on a LightCycler 480 II thermal cycler (Roche). Gene expression was normalized to *GAPDH*; see Table 2 for all primers.

Immunofluorescence staining

Cells were fixed using 4% PFA (20 min, RT), permeabilized with 0.1% Triton X-100 (Sigma-Aldrich; 15 min, RT) and blocked (45 min, RT) in 1% BSA (Sigma-Aldrich)/10% donkey serum (Millipore). Primary antibodies were incubated overnight, 4 °C (no primary antibody for negative control), and fluorescence-conjugated secondary antibodies 1 h, RT (Jackson ImmunoResearch). Nuclei were stained with 0.2 μ g/ml bisBenzimide (Sigma-Aldrich). Cells were observed using a Zeiss ApoTome 2 Upright wide-field microscope.

In vitro differentiation assay

iPSC were dissociated with Accutase (Stemcell Technologies) and seeded on ultra-low attachment dishes for 2 days in E8 containing

Table 2 Reagents details.

Antibodies used for immunocytochemistry/flow-cytometry

	Antibody	Dilution	Company Cat # and RRID
Pluripotency markers	Rabbit anti-SOX2	1/200	Thermo Fisher Scientific Cat# 48–1400, RRID:AB_2533841
Pluripotency markers	Rabbit anti-NANOG	1/200	Abcam Cat# ab21624, RRID:AB_446437
Pluripotency markers	Mouse anti-OCT3/4	1/200	Santa Cruz Biotechnology Cat# sc-5279, RRID:AB_628051
Differentiation markers	Rabbit anti-GFAP	1/200	Dako Cat# Z0334, RRID:AB_10013382
Differentiation markers	Mouse anti-SMA	1/200	Dako Cat# M0851, RRID:AB_2223500
Differentiation markers	Mouse anti-AFP	1/200	Sigma Aldrich Cat# WH0000174M1, RRID:AB_1839587
Secondary antibodies	Alexa Fluor® 488 AffiniPure Donkey Anti-Rabbit IgG (H + L)	1/500	Jackson ImmunoResearch, Cat# 711-545-152, RRID:AB_231358
Secondary antibodies	Alexa Fluor [®] 488 AffiniPure Donkey Anti-Mouse IgG (H + L)	1/500	Jackson ImmunoResearch, Cat# 715-545-150, RRID:AB 23408

Primers

	Target	Forward/Reverse primer (5'-3')	Product size (bp)	
Exogenous reprogramming transgene (RT-PCR)	SeV	GGATCACTAGGTGATATCGAGC	181	
		ACCAGACAAGAGTTTAAGAGATATGTATC		
Exogenous reprogramming transgene (RT-PCR)	KOS	ATGCACCGCTACGACGTGAGCGC	528	
		ACCTTGACAATCCTGATGTGG		
Exogenous reprogramming transgene (RT-PCR)	KLF4	TTCCTGCATGCCAGAGGAGCCC	410	
		AATGTATCGAAGGTGCTCAA		
Exogenous reprogramming transgene (RT-PCR)	c-MYC	TAACTGACTAGCAGGCTTGTCG	532	
		TCCACATACAGTCCTGGATGATGATG		
Pluripotency marker (RT-qPCR)	NANOG	CAAAGGCAAACAACCCACTT	158	
		TCTGCTGGAGGCTGAGGTAT		
Pluripotency marker (RT-qPCR)	OCT3/4	GTACTCCTCGGTCCCTTTCC	168	
		CAAAAACCCTGGCACAAACT		
Pluripotency marker (RT-qPCR)	LIN28a	GGGGAATCACCCTACAACCT	166	
		CTTGGCTCCATGAATCTGGT		
GAPDH RT-(qPCR and RT-PCR)	GAPDH	AACCATGAGAAGTATGACAAC	112	
		CTTCCACGATACCAAAGTT		
USH2A exon 13	USH2A	GAAGTTCATCGCAAACAGTTG	686	
		CACTGATTACAGCGAAGACCTG		

Y27632 StemMACS. At day 3, the medium was changed to DMEM/F12 (Gibco) supplemented with 20% Knockout serum replacement (Gibco), 1% penicillin-streptomycin (Gibco), 1% GlutaMax (Gibco), 1% NEAA (Gibco) and 55 mM β -mercaptoethanol (Gibco). At day 7, embryoid bodies were seeded onto Matrigel-coated wells and cultured in the same medium for a further 10 days before staining.

Mutation analysis

Genomic DNA was isolated using the DNeasy Blood & Tissue Kit (Qiagen) and PCR-amplified using *USH2A*-specific primers (Table 2). The dNTPs were removed using the ExoSAP-IT PCR Clean-up kit (GE Healthcare) and the amplicon sequenced using the BigDye Terminator Cycle Sequencing Ready Reaction kit V3.1 on an Applied Biosystems 3130xL Genetic Analyzer.

Microsatellite PCR analysis

Genomic DNA was amplified using primers for informative markers (Table 1). The PCR products were mixed with Genescan 400HD ROX size standard and subsequently analyzed on an Applied Biosystems 3130xL genetic analyzer.

Mycoplasma analysis

Mycoplasma detection was performed on cell culture supernatant using the MycoAlert Mycoplasma Detection Kit (Lonza), according to the manufacturer's instructions, and a CLARIOstar microplate reader (BMG Labtech).

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.scr.2018.11.007.

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