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# Generation of the human iPSC line ESi082-A from a patient with macular dystrophy associated to mutations in the *CRB1* gene

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Alberto Cañibano-Hernández<sup>a</sup>, Lourdes Valdes-Sanchez<sup>a</sup>, Ana B. Garcia-Delgado<sup>a</sup>, Beatriz Ponte-Zúñiga<sup>b</sup>, Francisco J. Diaz-Corrales<sup>a</sup>, Berta de la Cerda<sup>a,\*</sup>

<sup>a</sup> Department of Regeneration and Cell Therapy, Andalusian Molecular Biology and Regenerative Medicine Centre (CABIMER), Avda. Americo Vespucio, 24, 41092
Seville, Spain
<sup>b</sup> University Hospital Virgen Macarena, RETICS Oftared, Carlos III Institute of Health (Spain), Ministry of Health RD16/0008/0010, Seville, Spain

### ABSTRACT

Retinal dystrophies associated to mutations in the *CRB1* gene comprise a wide array of clinical presentations. A blood sample from a patient with a family history of *CRB1*-retinal dystrophy was used to prepare the iPSC line ESi082-A. The genotype of the donor, affected of a perifoveal-bilateral macular dystrophy includes one frameshift deletion and one hypomorphic allele. ESi082-A cell line has been characterized for pluripotency and will be used to prepare retinal cellular models to study the dysfunction leading to the disease.

### 1. Resource table

Unique stem cell line identifier	ESi082-A
Alternative name(s) of stem cell line	CRB1-MiPS4F1
Institution	Andalusian Molecular Biology and Regenerative
	Centre (CABIMER) Seville Spain
Contact information of	Berta de la Cerda
distributor	berta delacerda@cabimer es
Type of cell line	iPSC
Origin	Human
Additional origin info	Age: 56
	Sex: Male
	Ethnicity: European descent
Cell Source	Peripheral blood mononucleated cells
Clonality	Clonal
Method of reprogramming	hOCT3/4, hSOX2, hc-MYC, and hKLF4 via non-
	integrative Sendai virus
Genetic Modification	Yes
Type of Modification	Hereditary
Associated disease	Macular dystrophy
Gene/locus	CRB1/ Chr. 1: 197,268,204–197,478,455
	Mutations: c.613_619del (p.Ile205Aspfs*13)
	LOVD#CRB1_000175
	c.498_506del (p.Ile167_Gly169del) CinVar ID#96659
Method of modification	N/A
Name of transgene or	N/A
resistance	

### (continued)

Inducible/constitutive	N/A
system	
Date archived/stock date	10. 20. 2020
Cell line repository/bank	hpsreg.eu/cell-line/ESi082-A
Ethical approval	Cellular Reprogramming Board of Andalucía. Ethical
	Approval number: PR_01_2015

### 2. Resource utility

Most CRB1-associated retinal dystrophies present as Leber congenital amaurosis or retinitis pigmentosa but the hypomorphic allele CRB1 c.613\_619del has been associated to macular dystrophy. The iPSC line ESi082-A will be used to obtain retinal cellular models by differentiation, enabling the study of the mechanisms of disease leading to cellular degeneration.

### 3. Resource details

Retinal dystrophies associated to *CRB1* mutations cover a wide spectrum of clinical presentations and currently it is unclear which are the factors leading to the different degrees of disease (Khan et al., 2018; Quinn et al., 2017). Cellular modelling using patient-derived iPSCs is a promising tool to uncover the mechanisms of disease. A patient diagnosed with macular dystrophy associated to a composite homozygous genotype in the *CRB1* gene was the donor of the biological sample to

\* Corresponding author.

E-mail address: berta.delacerda@cabimer.es (B. de la Cerda).

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obtain the ESi082-A iPSC line. Mononucleated cells collected from peripheral blood (PBMCs) were reprogrammed to iPSCs using the nonintegrative Sendai virus to transduce the reprogramming factors hOCT3/4, hc-MYC, hKLF4, and hSOX2. Identity of the cell line was confirmed by DNA fingerprinting, matching with the DNA of the original biological material (available with the authors). Moreover, DNA sequencing of ESi082-A compared to a normal control confirmed the presence of the two mutations in the CRB1 gene (Fig. 1A). CRB1/ c.613\_619del mutation is the maternal allele, a mutation previously associated to Leber congenital amaurosis (http://databases.lovd.nl/sh ared/variants/CRB1), and CRB1/c.498\_506del is the paternal allele, reported as pathogenic for macular dystrophy (Khan et al., 2018). The iPSC line was checked for normal 46, XY male karyotype (Fig. 1B) and for the typical stem cell morphology of compact flat colonies composed of cells with a high nuclear/cytosolic ratio (Fig. 1D). Clearance of viral capsid and ectopic reprogramming factors was apparent at passage 7, with no expression of these genes detected by RT-PCR (Fig. 1C). To characterize the pluripotency of cell line, expression of *NANOG* and *OCT4* genes was quantified by RT-qPCR. ESi082-A iPSC line presented a high level of expression for both genes compared to PBMCs of the patient, used as a negative control (Fig. 1G). A complete panel of pluripotency markers was evaluated by immunofluorescence, demonstrating nuclear expression for OCT4, NANOG and SOX2, and surface expression for SSEA3, SSEA4 and TRA-1-81 (Fig. 1F; right panels depict a magnification of each left panel). Additionally, elevated enzymatic activity for alkaline phosphatase was also shown for the iPSC line (Fig. 1E). The

### Table 1

Characterization and validation.

Classification	Test	Result	Data
Morphology Phenotype	Photography Qualitative analysis	Normal morphology Positive staining for pluripotency markers: OCT4, NANOG, SOX2, TRA-1-81, SSEA3, SSEA4 and alkaline phosphatase.	Fig. 1D Fig. 1F, 1E
	Quantitative analysis	RT-qPCR analysis of the expression for NANOG and OCT4	Fig. 1G
Genotype	Karyotype (G- banding) and resolution	46, XY, Resolution 450–500	Fig. 1B
Identity	Microsatellite PCR (mPCR) OR STR analysis	STR analysis. 10 sites tested. All matched mPCR not done	Available with the authors N/A
Mutation analysis	Sequencing	chr1:197,201,504: c.613_619del c.498_506del	Fig. 1A
	Southern Blot OR WGS	Not done	N/A
Microbiology and virology	Mycoplasma	Negative <i>Mycoplasma</i> detection. Tested by luminescence	Supplementary Fig. 1
Differentiation potential	In vitro embryoid body formation	Positive immunostaining for mesodermal muscle actin, endodermal alpha fetoprotein and ectodermal PAX6	Fig. 1H
Donor screening	HIV 1 + 2 Hepatitis B, Hepatitis C	Not done	N/A
Genotype additional info	Blood group genotyping	Not done	N/A

ability of ESi082-A to generate the three germ layers was evaluated by inducing the formation of embryoid bodies *in vitro* and then confirming the expression of specific markers by immunofluorescence: endodermal  $\alpha$ -fetoprotein (AFP), mesodermal smooth-muscle actin (SMA) and ectodermal PAX6 (Fig. 1H; right panels depict a magnification of each left panel). Characterization is summarized in Table 1.

### 4. Materials and methods

### 4.1. Mutation sequencing

Genomic DNA from iPSCs was isolated using the QIAamp DNA Blood mini kit (Qiagen). Primers for amplification and Sanger sequencing of the genomic region flanking the mutation sites are described in Table 2.

### 4.2. Peripheral blood mononuclear cells (PBMCs) primary culture

PBMCs were isolated from 4 mL of peripheral blood using Vacutainer CPT tubes (BD) and cultured in expansion medium (EM: QBSF-60 (Quality Biological) with 50  $\mu$ g/ml of ascorbic acid (Sigma-Aldrich); 50 ng/ml of SCF (Stemcell Technologies); 10 ng/ml IL-3 (Stemcell Technologies); 2 U/mL of EPO (Stemcell Technologies); 40 ng/ml IGF-1 (Stemcell Technologies); 1  $\mu$ M Dexamethasone (Sigma-Aldrich) and 1% Pen/strep (Gibco)).

### 4.3. Reprogramming PBMCs to iPSCs

 $2.5 \times 10^4$  PBMCs were transduced with reprogramming factors (CytoTune®-IPS Reprogramming Kit (Life Technologies)) for 24 h. Cells were washed and kept in EM medium for 48 h.  $1 \times 10^5$  cells were seeded

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Table 2	
Reagents	details.

Antibodies used fo	or immunocytoche Antibody	mistry/flow-cy Dilution	company Cat # and RRID	
Pluripotency Markers	Rabbit anti- OCT4	1:400	Cell Signaling Technology	
Warkers	Rabbit anti-	1.400	Cell Signaling Technology	
	NANOG	1.100	Cat# 4903_BBID	
	inniou		AB 10559205	
	Rabbit anti-	1:400	Cell Signaling Technology	
	SOX2		Cat# 3579, RRID:AB 2195767	
	Mouse anti-	1:100	Millipore Cat# MAB 4304,	
	SSEA4		RRID: AB_177629	
	Mouse anti-	1:100	Stemgent Cat# 09–0069, RRID	
	TRA-1-81		AB_2119069	
	Rat anti-	1:100	Thermo Fisher Cat# MA 1–020	
	SSEA3		RRID: AB_2536682	
Differentiation	Rabbit anti-	1:500	Covance Cat# PRB-278P,	
Markers	PAX6		RRID: AB_291612	
	Mouse anti-	1:300	Sigma-Aldrich Cat# A5228,	
	SMA		RRID: AB_262054	
	Mouse anti-	1:20	R & D Systems Cat# MAB1368	
	AFP		RRID: AB_357658	
Secondary	Donkey anti-	1:500	Thermo Fisher Scientific Cat#	
antibodies	Rabbit 594	1 500	A-21207, RRID: AB_141637	
	Donkey anti-	1:500	Thermo Fisher Scientific Cat#	
	Mouse 488	1.500	A-21202, RRID: AB_141607	
	Rabbit anti-	1:500	Invitrogen Cat# A21210,	
During and	Nat		RRID. RD_2000700	
Primers	Torgot size	Forward /Box	(E', 2')	
	(bp)	FOI WAI U/ Kev	verse primer (3 -3 )	
SeV genome	SeV, 181	GGATCACTAGGTGATATCGAGC/		
silencing (RT- PCR)		ACCAGACAA	AGAGTTTAAGAGATATGTATC	
	KOS, 528	ATGCACCGCTACGACGTGAGCGC/		
		ACCTTGACAATCCTGATGTGG		
	KLF4, 410	TTCCTGCATGCCAGAGGAGCCC/		
		AATGTATCGAAGGTGCTCAA		
	cMYC, 532	TAACTGACTAGCAGGCTTGTCG/		
		TCCACATACAGTCCTGGATGATGATG		
House-Keeping	GAPDH, 86	ACGACCCCTTCATTGACCTCAACT/		
Gene (RT- PCR)		ATATTTCTCGTGGTTCACACCCAT		
Pluripotency markers (RT- aPCR)	NANOG	Hs02387400	g1	
4	OCT4	Hs01654807	s1	
House-keeping gene (RT-	GADPH	Hs02786624g1		
(PCK)	A11010	CCAACAAC	CCTCCCAAC/	
l'argeted	Allele	GCAAGAACI	ILLIGUCAAC/	
analysis/	del, 262	AGGCAIGTA	4966169116	
sequencing	Allele	TACTCCTGC	TTCTGTGTCC/	
	c.498–506 del. 170	GTGATTCAG	GATATGATCACACTTACC	

on feeders (irradiated hFF (ATCC CRL2429)) and cultured in QBSF-60 with 50  $\mu$ g/ml of ascorbic acid and 1% pen/strep. On day 15, medium was changed to hESCM: Knock Out DMEM-F12 (Gibco); 20% Knock Out serum (Gibco); 1% glutamine (Gibco); 1% NEAA (Gibco); 0.23 mM  $\beta$ -mercaptoethanol (Gibco); 1% pen-strep and 10 ng/ $\mu$ l bFGF (Peprotech)). Colonies with stem-like morphology were manually isolated in the days 21 to 27 post-transduction. Clone 1, named ESi082-A, was adapted to grow in mTeSR1 (Stemcell Technologies) onto Matrigel.

## 4.4. RT-PCR and qPCR for detection of viral clearance and pluripotency markers

 $1 \ \mu g$  of total RNA isolated with RNeasy Mini Kit (Qiagen) was retrotranscribed using QuantiTect Reverse Transcription Kit (Qiagen). RT- PCR reaction was performed using MyTaq DNA Polymerase (Bioline GmbH) with the primers listed in Table 2. For Q-PCR, triplicate reactions were prepared with TaqMan Gene expression Master Mix and probes listed in Table 2.

### 4.5. Immunofluorescence analysis

Cells grown in glass coverslips were washed with PBS, fixed in 4% PFA for 15 min, washed twice in PBS and blocked with 2% donkey serum in TPBS (0.2% Triton-X100/PBS) for 1 h. Primary antibody was incubated for 1 h in 1% BSA in TPBS, washed 3 times in TPBS and incubated with secondary antibody diluted in 1% BSA in TPBS for 1 h. After 3 washes, preparations were finished with mounting medium with DAPI. Antibodies are listed in Table 2.

### 4.6. Alkaline phosphatase

Enzymatic activity was detected using AP staining kit II (Stemgent).

### 4.7. In vitro differentiation

iPSC colonies were manually collected and cultured in non-adherent conditions in hESCM without bFGF for 7 days. Embryoid bodies were manually collected and seeded onto gelatinized-glass coverslips, cultured for one week in EBM (DMEM/F12, 10% FBS, 1% GlutaMAX, 1% MEM NEAA, 1% Pen/Strep) and analyzed by immunofluorescence using the antibodies in Table 2.

### 4.8. Karyotype analysis

On passage 11, 30 metaphases were counted and genome integrity was analyzed by G-banding at 400–550 band resolution in Biobanco del SSPA, Granada, Spain.

### 4.9. Fingerprinting

STR analysis was performed form iPSC and PBMC genomic DNA

using the GenePrint  $\ensuremath{\mathbb{R}}$  10 System (Promega) in Biobanco del SSPA, Granada, Spain.

### 4.10. Mycoplasma detection

MycoAlert<sup>™</sup> PLUS *Mycoplasma* Detection Kit (Lonza) was used to test for *Mycoplasma* (Suppl. Fig. 1).

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### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

### Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.scr.2021.102301.

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