

Invoice number: SCTC2018-00081

Certificate of Analysis 2019

Name investigator: Rob Collin Cell line number: IPS18-00092

Project name: FFB

Table 1: Information on the reprogrammed cell line

Information cell line:	
Product description	PBMCs nucleofected with episomal vectors containing the genes OCT3/4, SOX2, KLF4, L-MYC, LIN28 PBMCs
Parental cell line	HEP18-00190
Parental cell type	PBMCs
Diagnosis	LCA
Mutation	
Number of clones	3
Passage (P) of iPS cells reported at submission	P6
Culture medium	Essential 8 Flex medium
Culture coating	Matrigel
Feeders during reprogramming	Mouse Embryonic Fibroblasts (MEFs)
Passage method	0.5 mM EDTA
Protocols in Q-portal	046588; 046591
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Table 2: Information on the characterization of the reprogrammed cell line

Test description:	Test method:	Test specification:	Result:
Activation of stem cell markers	qPCR	Upregulation of <i>OCT4, SOX2, NANOG, DNMT3B</i> compared with PBMCs	Pass
Expression of stem cell markers	Immunocytochemistry	Expression of OCT4, NANOG, SSEA4	Pass
Mycoplasma	PCR	Negative	Pass
Three lineage	Differentiation assay	Upregulation of germlayer specific	N/A*
differentiation		genes	

^{*}N/A: Not Applicable

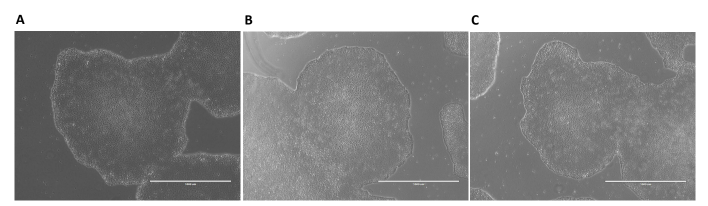


Figure 1: Cells prior to freezing. A - C, clone 1, 2 and 3, respectively at P6. Scale bar = 1000 μm .





All clones were assessed for activation of stem cell markers before freezing. RNA was isolated and gene expression was assessed by quantitative reverse transcription PCR. Ct values were normalized with the housekeeping gene GUSB (set at 1).

Absolute expression, normalized to GusB

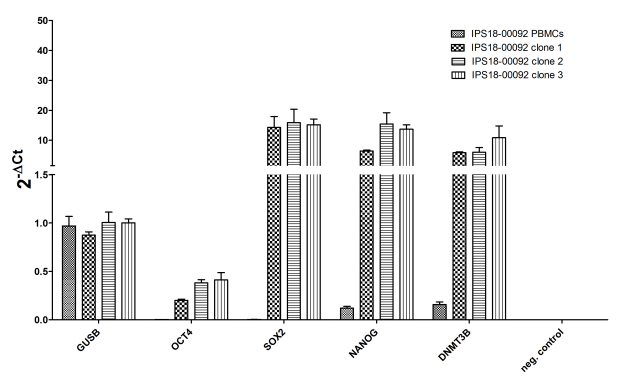


Figure 2: Gene expression of three iPS cell clones compared with the parental PBMCs (ΔCt).

Expression relative to parental line

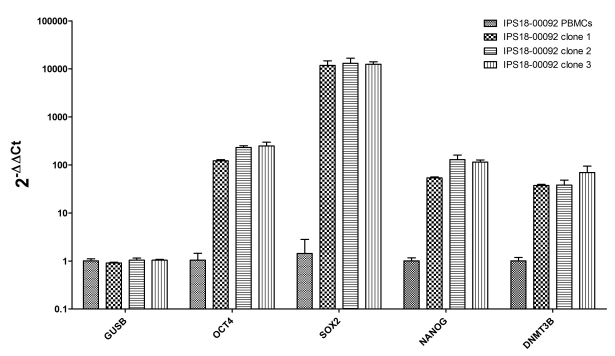


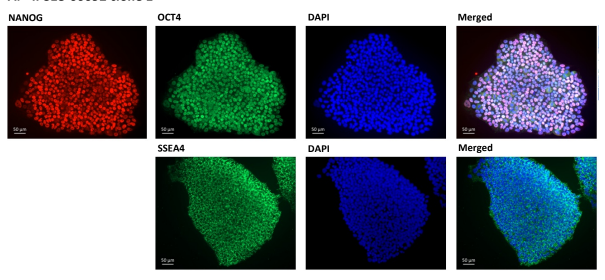
Figure 3: Pluripotency gene upregulation after reprogramming ($\Delta\Delta$ Ct). The expression fold difference of the iPS clones is relative to the parental PBMCs.



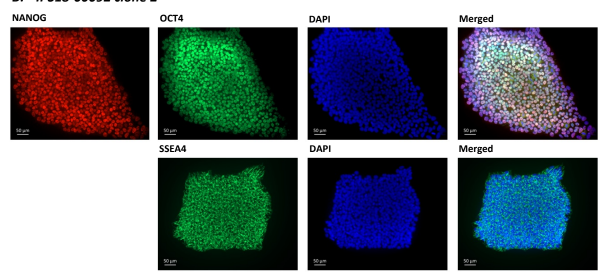
Expression of stem cell markers

Undifferentiated iPS cell clones were stained for the nuclear markers NANOG and OCT4, and surface antigen SSEA4. All markers are expressed in human pluripotent stem cells.

A. IPS18-00092 clone 1



B. IPS18-00092 clone 2





C. IPS18-00092 clone 3

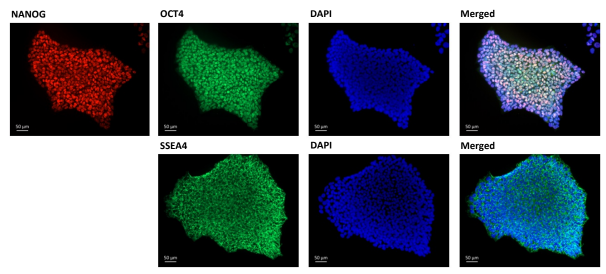


Figure 4. Immunofluorescence staining of the 3 iPS cell clones (A, B, C) with pluripotency markers.

Pass

Fail

Other:

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