

Invoice number: SCTC2021-00074

Certificate of Analysis 2021

Name principal investigator: Frans Cremers Cell line number: IPS15-00006 Project name: -

Table 1: Information on the reprogrammed cell line

Information cell line:	
Product description	Human fibroblasts reprogrammed with four factors (OCT4, SOX2, KLF4, C-MYC) by using lentiviral vectors
Parental cell line	CL14-00047
Parental cell type	Fibroblasts
Diagnosis	STGD1
Mutation	N/A*
Number of clones	3
Passage (P) of iPSCs reported at delivery	P6
Culture medium	Essential 8 Flex medium
Culture coating	Vitronectin
Feeders during reprogramming	Mouse Embryonic Fibroblasts (MEFs)
Passage method	EDTA

*N/A: Not Applicable

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 OCT4, SOX2, LIN28, NANOG in iPSCs compared with PBMCs	Pass
Expression of stem cell markers	Immunocytochemistry	Expression of OCT4, NANOG, SSEA4, TRA-1-81	Pass
Mycoplasma test	PCR	Negative	Pass
Three lineage differentiation	Differentiation assay	Upregulaton of germlayer-specific markers	Pass

Α



В





Figure 1: Cells prior to freezing. A - C, clones 1 - 3 at P6.

Activation of stem cell markers

The RNA of all clones was isolated before freezing and the gene expression was assessed by quantitative reverse transcription PCR (qRT-PCR). Ct values were normalized with the housekeeping gene GUSB, set at 1.

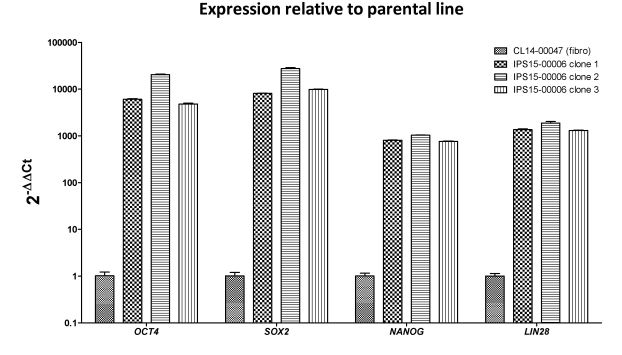
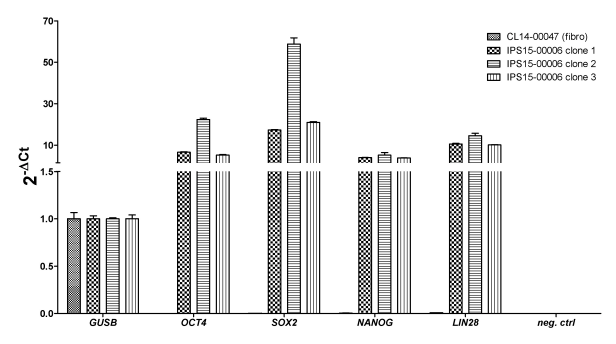


Figure 2: Gene expression of three iPSC clones compared with the parental fibroblasts (Δ Ct).

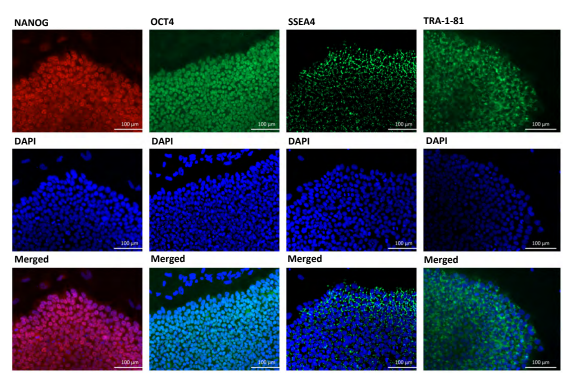


Absolute expression, normalized to GusB

Figure 3: Pluripotency gene upregulation after reprogramming (ΔΔCt). The expression fold difference of the iPSCs is relative to the parental fibroblasts.

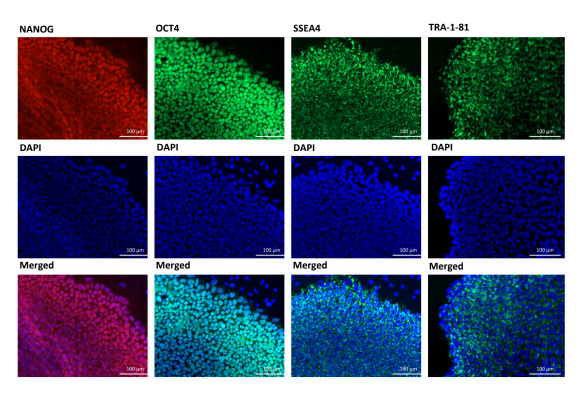
Expression of stem cell markers

The undifferentiated iPSC clones were stained for the nuclear markers NANOG and OCT4 and surface antigens SSEA4 and TRA-1-81. All markers are expressed in human pluripotent stem cells.



A. IPS15-00006 clone 1

B. IPS15-00006 clone 2



C. IPS15-00006 clone 3

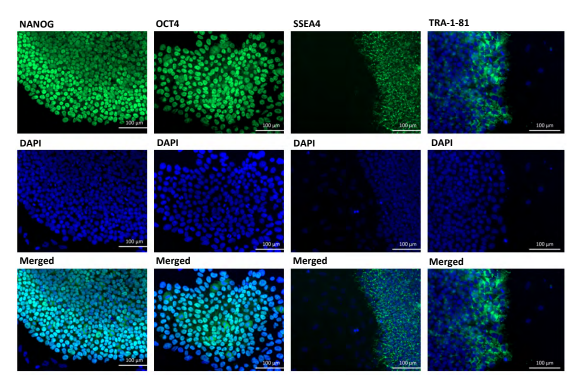


Figure 4: Immunofluorescence staining of the iPSC clones with pluripotency markers.

Three germ layer differentiation

Clone 1 at passage 12 was differentiated into the endodermal, mesodermal and ectodermal germ layers. The RNA was isolated and the gene expression was checked by qPCR. Ct values are normalized with the housekeeping gene GUSB, set at 1. For each lineage two genes were assessed (Table 3). The differentiated cells were also stained for lineage-specific markers (Table 4).

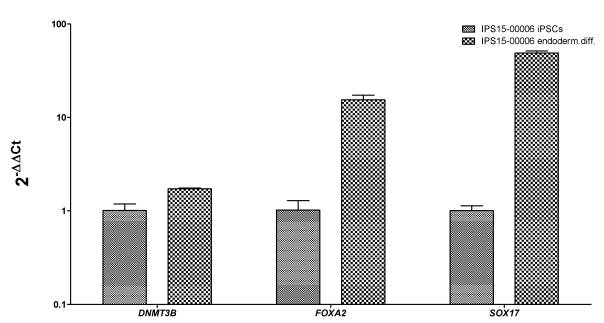
Table 3: qPCR markers for three lineage differentiation

Lineage	Marker
Endoderm	FOXA2, SOX17
Mesoderm	Brachyury, HAND1
Ectoderm	PAX6, NCAM1

Table 4: ICC markers for three lineage differentiation

Lineage	Marker
Endoderm	SOX17
Mesoderm	DESMIN
Ectoderm	NESTIN

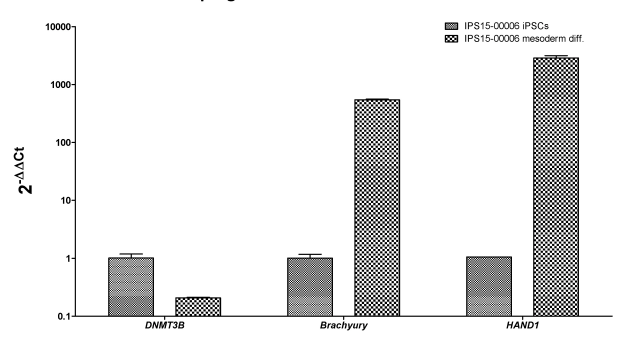
Endoderm



Upregulation of endodermal markers

Figure 5: Expression fold difference of endoderm-specific genes in differentiated cells, compared with undifferentiated iPSCs. *DNMT3B* was used as a reference for pluripotency.

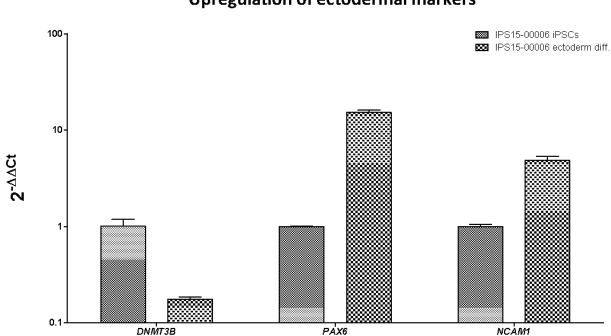
Mesoderm



Upregulation of mesodermal markers

Figure 6: Expression fold difference of mesoderm-specific genes in differentiated cells, compared with undifferentiated iPSCs. *DNMT3B* was used as a reference for pluripotency.

Ectoderm



Upregulation of ectodermal markers

Figure 7: Expression fold difference of ectoderm-specific genes in differentiated cells, compared with undifferentiated iPSCs. *DNMT3B* was used as a reference for pluripotency.

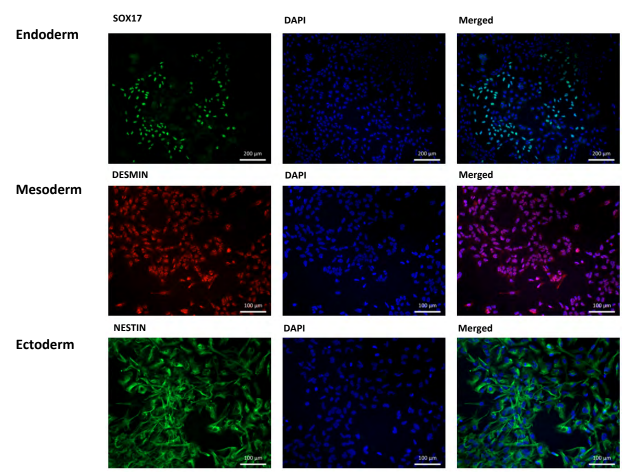


Figure 8: Immunofluorescence staining of differentiated cells showing a positive signal of germlayer-specific markers.

Pass

Fail

Other:

Silvialbes

Silvia Albert, PhD Manager, Radboud Stem Cell Technology Center Date