

Certificate of analysis

SFC049-03-13

Signature: Theodore Latsis: 11-08-2015

Supervisor signature: Lyle Armstrong

Date: 11-08-2015

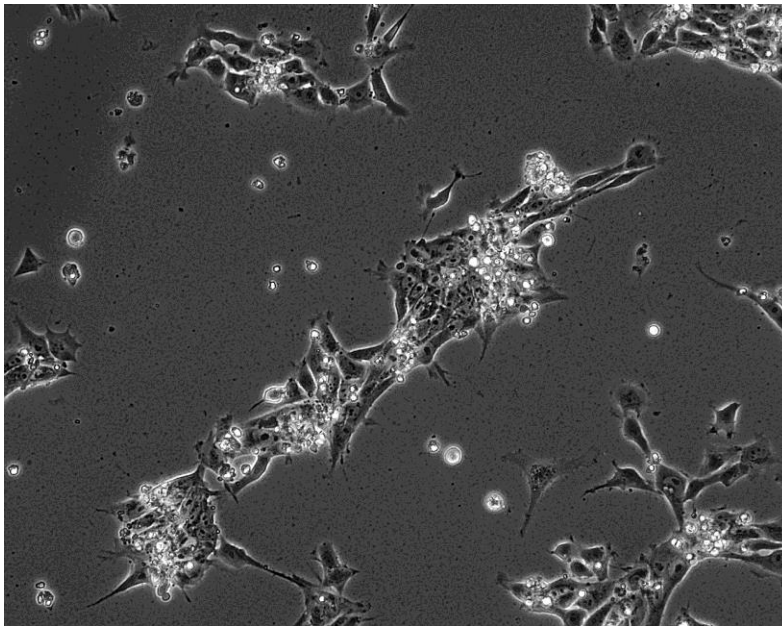
Source of fibroblasts and reprogramming information

- SF049 from University of Oxford
- Reprogrammed at UNEW, on 13-01-2015 at passage 5
- Cytotune 2
- This cell line has been difficult to expand mainly due to differentiation when passaged. Over time, this clone became accustomed to EDTA (versene) passaging and differentiation was diminished. Cells/colonies should be passaged every 4-6 days. Cultures should not be allowed to become confluent as it will lead to differentiation.

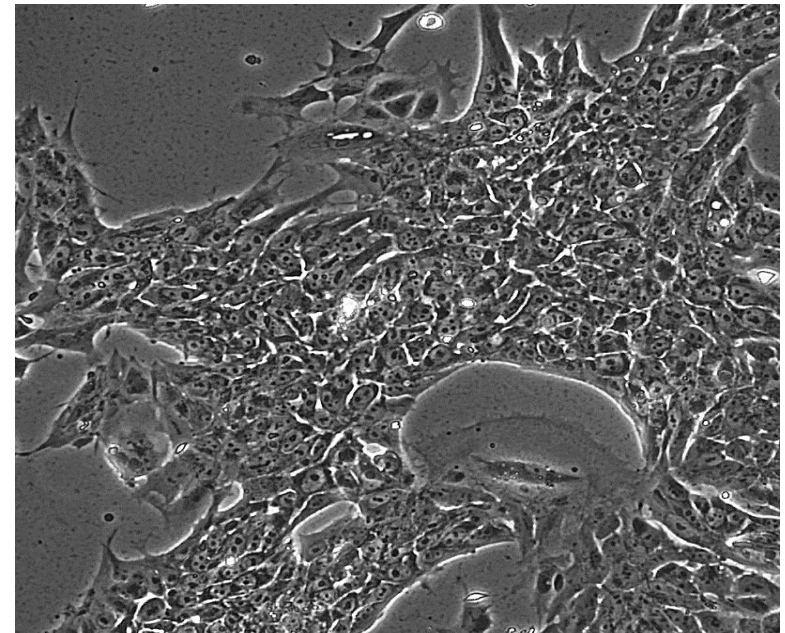
Viability post-thaw and Morphology according to SOP19 passage 20

- Cell count immediately post-thaw: 1.5×10^6
- Viability immediately post-thaw: 87%
- Photo 24 & 32h post-thaw

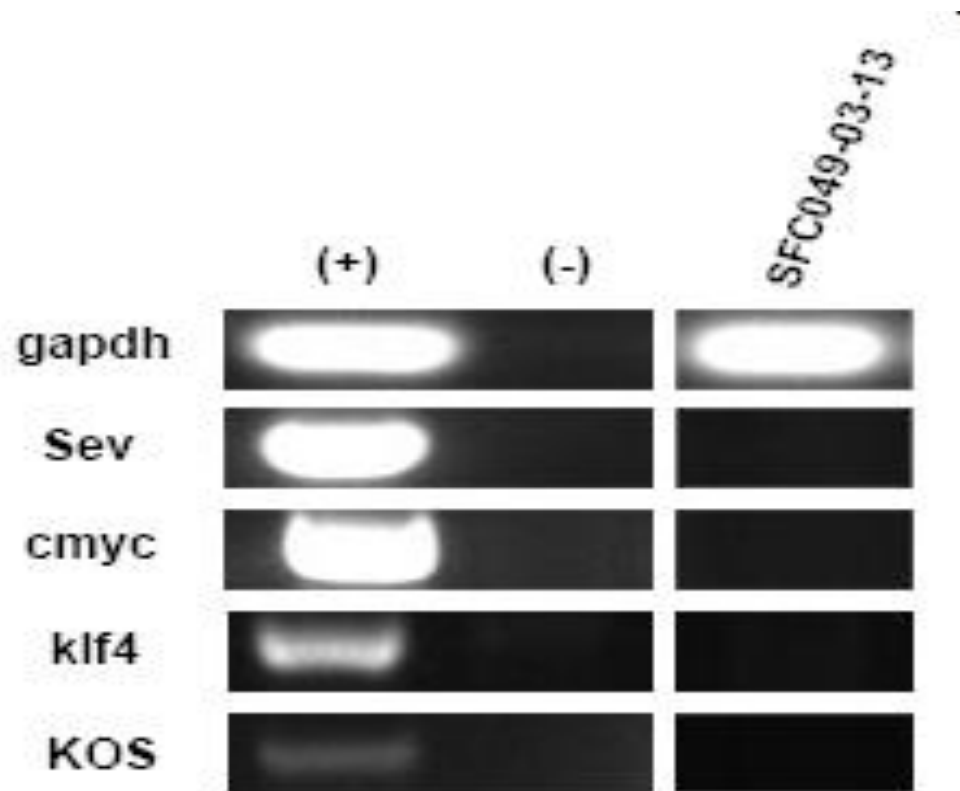
24h (x10)



32h (x10)



Sendai clearance:
according to WP3 SOP15
undetectable at passage 20



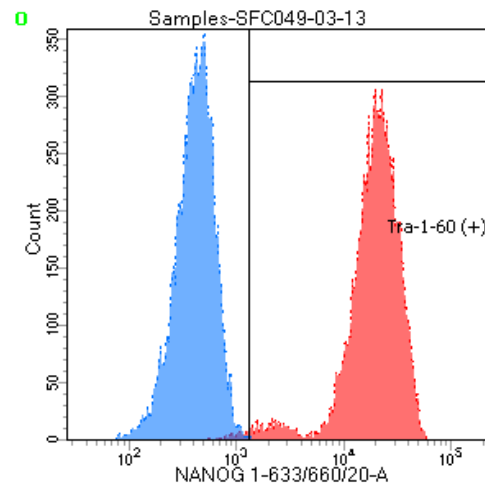
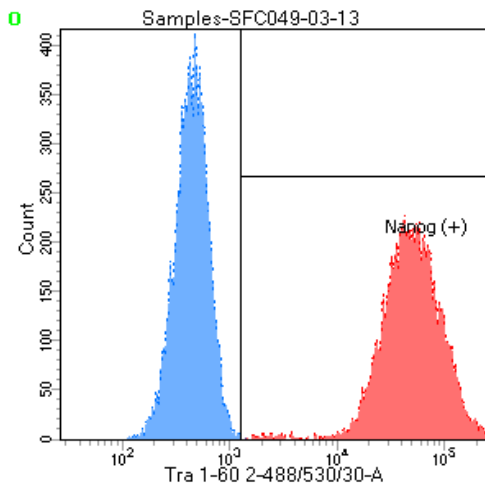
Mycoplasma test:

According to MycoAlert Lonza LT07-318

Undetectable at passage 20

> 1.2		Mycoplasma Contaminated	Positive Control	Negative Control	Cell name	SFC-049-03-13
0.9-1.2		Status Unknown - Restest within 24 hours	0.053	0.062	A	0.018
0-0.9		Mycoplasma Free	0.963	0.010	B	0.010
			18.068	0.166	B/A	0.549

Flow cytometric analysis according to WP3 SOP 20 and 21 passage 20



Tube: SFC049-03-13

Population	#Events	%Parent	%Total
■ All Events	20,000	####	100.0
■ Cells	9,339	46.7	46.7
☒ Tra-1-60 (+)	9,224	98.8	46.1
☒ Nanog (+)	9,334	99.9	46.7
■ Samples/49/All Events	20,000	####	100.0
■ Samples/49/P1	9,231	46.2	46.2

SNP analysis

according to WP3 SOP Preparation of DNA and RNA samples for Illumina arrays

- Passage 20
- Identity to parent fibroblasts confirmed
- Karyotype abnormalities: none detected
- For details and raw data see StemDB
- Audit 21.07.17