

Certificate of analysis

SFC888-07-02

Signature: Dario Melguizo Sanchis

Date: 12.06.2015

Supervisor signature: Linda Lako

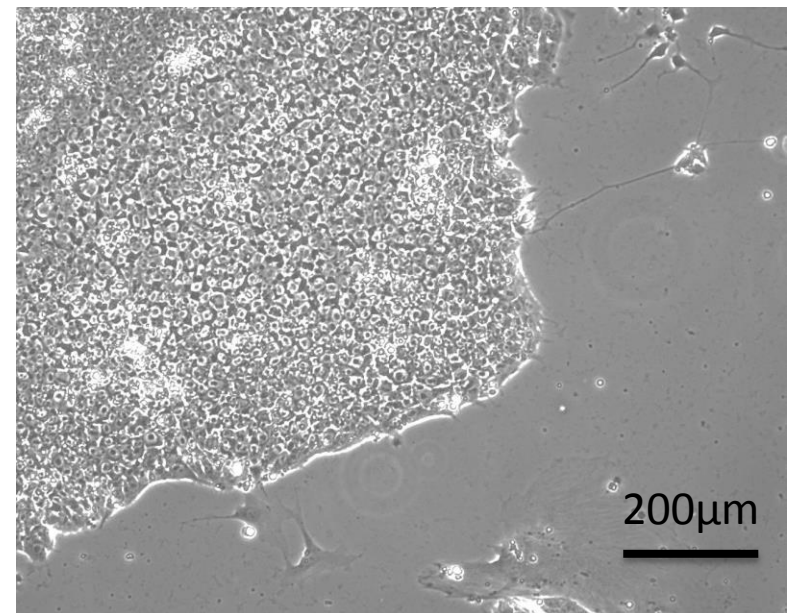
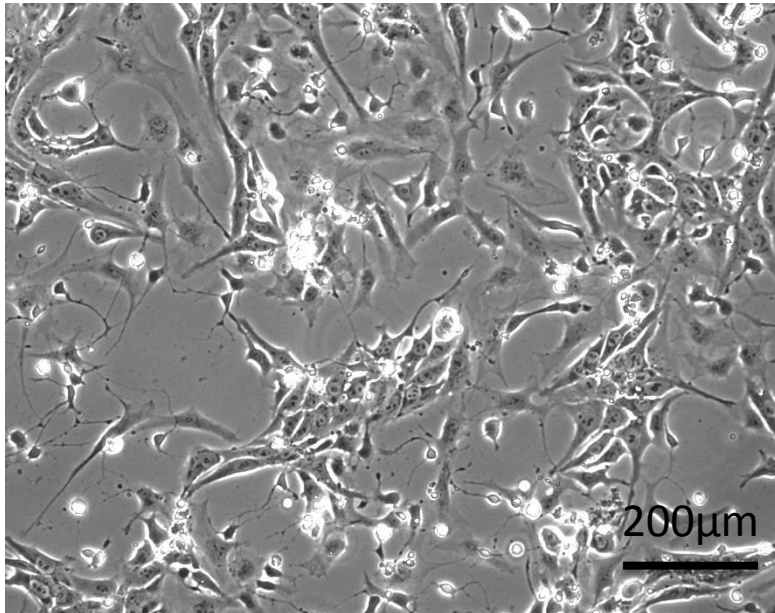
Date: 12.06.2015

Source of fibroblasts and reprogramming information

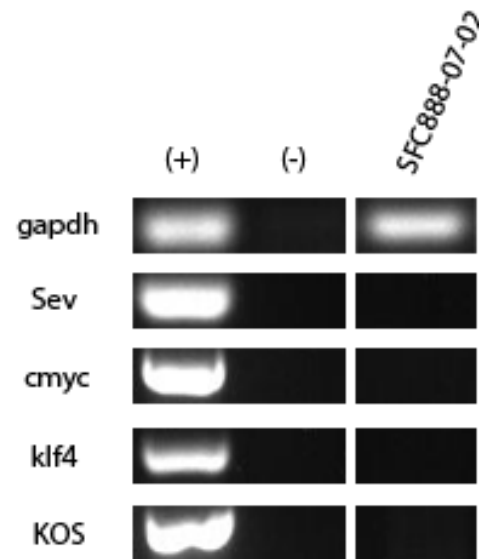
- SF888 from UOXF 01/15
- Reprogrammed at UNEW ISV
- Reprogrammed on 17/02/2015 at passage 8
- Cytotune v2 WP3 SOP22

Viability post-thaw and Morphology according to SOP19 passage 10

- Cell count immediately post-thaw 2×10^6
- Viability immediately post-thaw 85%
- Photo at 24h post thaw (left) and 3d after ReLSR passaging (right):



Sendai clearance:
according to WP3 SOP15
undetectable at passage 10



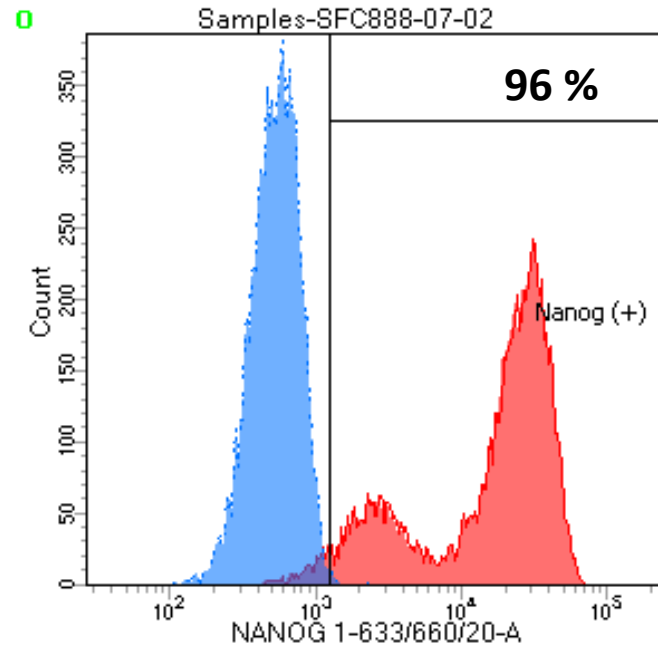
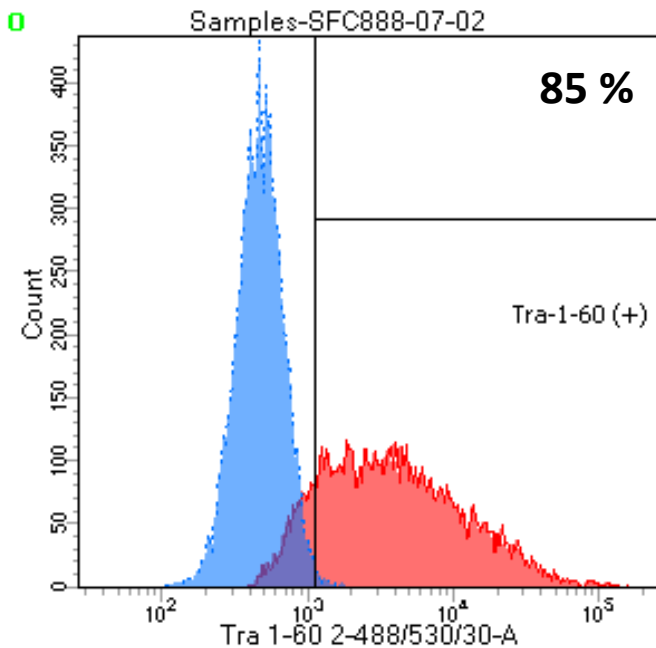
Mycoplasma test: Undetectable at passage 10

Owner	DM					
Date	08/06/2015					
Cell name	SFC888-07-02					
A	0.0264					
B	0.0119					
B/A	0.450757576					
> 1.2		Mycoplasma Contaminated			Positive Control	Negative Control
0.9-1.2		Status Unknown - Restest within 24 hours			0.0231	0.0862
0-0.9		Mycoplasma Free			1.769	0.0062
					76.58008658	0.071925754

Flow cytometric analysis according to WP3 SOP 20 and 21 passage 10

Tra-1-60:

NANOG:



SNP analysis

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

- Passage 10
- Identity to parent fibroblasts confirmed
- Karyotype abnormalities: none detected
- For details and raw data see StemDB

Comments

- Thawing in 2 wells of 6 well-plate recommended
- Clone prone to differentiation (see flow results weak positive population (2nd peak)) > ReLESR
Passaging after thawing recommended
 - Add 1ml ReLESR, incubate for 3 min at 37°C
 - Remove ReLESR
 - Add medium **dropwise** to collect undifferentiated cells