

### **Information Summary**

**Organization:** Neural Stem Cell Institute (NSCI) **Address:** One Discovery Drive, Rensselaer, NY 12144

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Phone: 518-694-8188 x 226

SCRC Code: NSCI11
Sample Name: FTD-T

Number of the Established iPSC Clones: 6

#### **Characterization Performed:**

1. Mycoplasma Test

2. Cell authentication Verification

3. Presence of the Pluripotency Marker

4. Spontaneous differentiation via embryoid body (EB) formation

5. Karyotyping

Qualified iPSC Clones: C1, C2, C4



## **Mycoplasma Test**

Method: Determination of the mycoplasmal enzyme activity using Lonza MycoAlert Plus Detection kit.

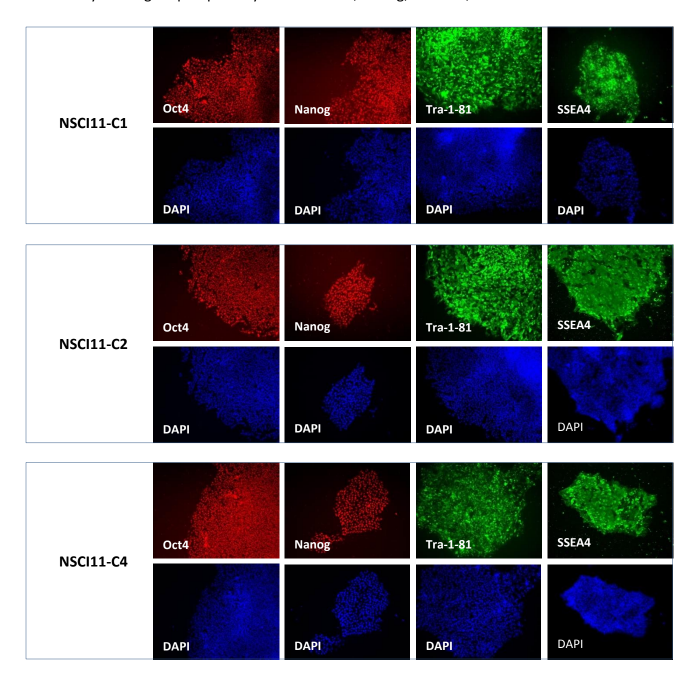
iPSC Clone	NSCI11-C1	NSCI11-C2	NSCI11-C4
Cell passage	P7	Р6	P6
B/A Ratio	0.53	0.24	0.54
Mycoplasma	NEG	NEG	NEG
Pass or Fail	Pass	Pass	Pass

## **Cell Authentication Verification**

Markers	NSCI11 (FTD-T)	NSCI11-C1	NSCI11-C2	NSCI11-C4
D3S1358	15, 18	15, 18	15, 18	15, 18
TH01	7, 9	7, 9	7, 9	7, 9
D21S11	27, 28	27, 28	27, 28	27, 28
D18S51	12, 16	12, 16	12, 16	12, 16
Penta E	8, 11	8, 11	8, 11	8, 11
D5S818	10, 12	10, 12	10, 12	10, 12
D13S317	11, 12	11, 12	11, 12	11, 12
D7S820	10, 13	10, 13	10, 13	10, 13
D16S539	11, 12	11, 12	11, 12	11, 12
CSF1PO	10	10	10	10
Penta D	11, 13	11, 13	11, 13	11, 13
vWA	17, 19	17, 19	17, 19	17, 19
D8S1179	10, 15	10, 15	10, 15	10, 15
TPOX	8, 11	8, 11	8, 11	8, 11
FGA	22, 23	22, 23	22, 23	22, 23
AMEL	X, Y	X, Y	X, Y	X, Y
Match with original Fibroblast		Yes	Yes	Yes

## **Expression of the Pluripotency Markers**

Antibody staining for pluripotency markers: Oct4, Nanog, Tra-1-81, SSEA-4





#### **Differentiation Potential via EB Formation**

**Method**: The iPSC cells were disassociated into small clumps to form EB in the low attachment dish. After a 10-day spontaneous differentiation, the EBs were harvested and RNAs were extracted. The expression levels of the selected 3-germ layer specific genes were analyzed by real-time PCR. Ct values are normalized for loading using a housekeeping gene.

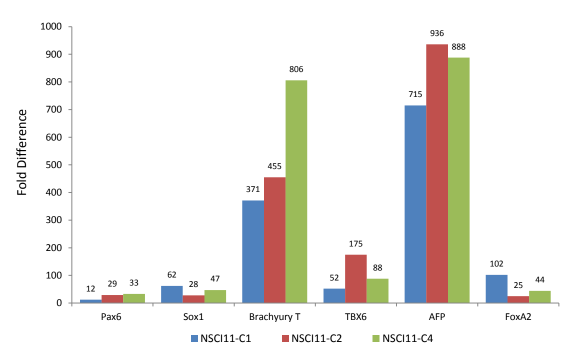


Figure: Lineage specific gene expression following EB differentiation. Fold difference is shown relative to undifferentiated iPS cell.



## Karyotyping

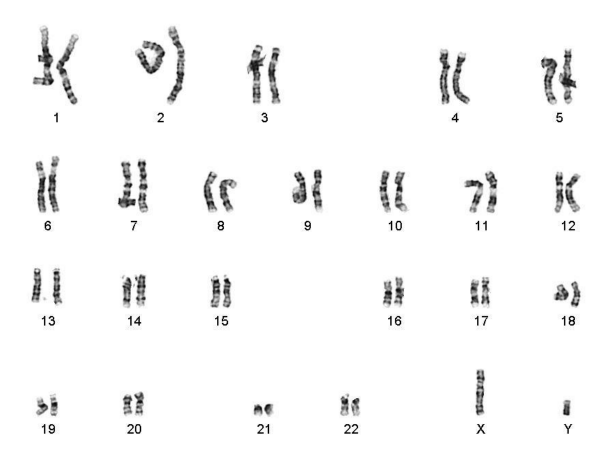
Clone: NSCI11-C1

Passage: P7

**Tested:** Number of metaphases counted: 20 Banding and level: GTG/550

Number of metaphases analyzed: 6 Number of karyotypes: 2

**Result:** 46,XY NORMAL MALE KARYOTYPE





### Karyotyping

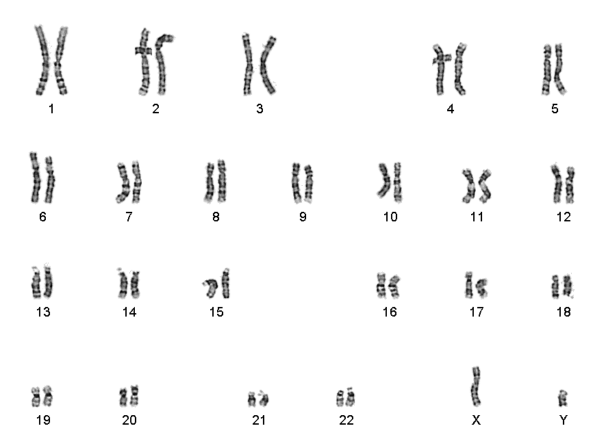
Clone: NSCI11-C2

Passage: P7

**Tested:** Number of metaphases counted: 20 Banding and level: GTG/450

Number of metaphases analyzed: 6 Number of karyotypes: 2

**Result:** 46,XY NORMAL MALE KARYOTYPE





## Karyotyping

Clone: NSCI11-C4

Passage: P6

**Tested:** Number of metaphases counted: 20 Banding and level: GTG/550

Number of metaphases analyzed: 6 Number of karyotypes: 2

**Result:** 46,XY NORMAL MALE KARYOTYPE

