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Transient p53 Suppression Increases Reprogramming of Human Fibroblasts without Affecting Apoptosis and DNA Damage

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SUMMARY

The discovery of human-induced pluripotent stem cells (iPSCs) has sparked great interest in the potential treatment of patients with their own in vitro differentiated cells. Recently, knockout of the Tumor Protein 53 (p53) gene was reported to facilitate reprogramming but unfortunately also led to genomic instability. Here, we report that transient suppression of p53 during nonintegrative reprogramming of human fibroblasts leads to a significant increase in expression of pluripotency markers and overall number of iPSC colonies, due to downstream suppression of p21, without affecting apoptosis and DNA damage. Stable iPSC lines generated with or without p53 suppression showed comparable expression of pluripotency markers and methylation patterns, displayed normal karyotypes, contained between 0 and 5 genomic copy number variations and produced functional neurons in vitro. In conclusion, transient p53 suppression increases reprogramming efficiency without affecting genomic stability, rendering the method suitable for in vitro mechanistic studies with the possibility for future clinical translation.

INTRODUCTION

Dr. Wilmut and colleague’s discovery in 1996 that mammalian somatic cells can be reprogrammed into a totipotent state of development by fusion with an enucleated oocyte (Campbell et al., 1996) has paved the way for autologous stem cell therapy by exploiting a patient’s own in vitro differentiated cells. Ten years later, with the revolutionizing finding that induced pluripotent stem cells (iPSCs) can be established from somatic cells by overexpression of merely four transcription factors (Takahashi and Yamanaka, 2006), a major step toward a clinical translation was taken. However, new coding mutations were recently reported to arise during the reprogramming process (Gore et al., 2011; Laurent et al., 2011; Martín-Taylor and Xu, 2012; Ng et al., 2011), rendering the process too hazardous for regenerative use in humans.

Several nonintegrative reprogramming strategies including Sendai virus (Fusaki et al., 2009) and episomal plasmids (Okita et al., 2011, 2013; Yu et al., 2009) have been tested to produce safe, transplantable iPSCs. Yet, recent whole-genome sequencing studies have failed to demonstrate a reduction in the mutational load in iPSC lines derived by integration-free reprogramming methods (Gore et al., 2011).

The cell-cycle regulator p53 acts as an important safeguard mechanism, by preventing cells from undergoing uncontrolled proliferation in response to DNA damage (Hong et al., 2009). The downstream DNA damage response (DDR) involves a series of events that lead to either cell-cycle arrest induced by p21 or apoptosis induced by PUMA. Importantly, p53 has also been shown to act as a critical barrier to the reprogramming process, and knockout of the TP53 gene in mouse and human fibroblasts was shown to produce significantly more iPSC colonies (Hong et al., 2009; Kawamura et al., 2009; Martín et al., 2009). Although the inhibition of p53 is clearly advantageous for the reprogramming efficiency, knocking out TP53 may possess critical safety issues because it was found to cause genomic instability (Chen et al., 2012; Lake et al., 2012; Martín et al., 2009; Menendez et al., 2010).

Recently, transient suppression of p53 with nonintegrative plasmids was shown to improve the reprogramming efficiency of human fibroblasts (Okita et al., 2011) and blood cells (Okita et al., 2013) by use of nonintegrative plasmids. Yet it remains unknown whether transient suppression of p53 will also give rise to genomic instability, which may in turn have detrimental effects on the gene expression, the epigenetic status, and the differentiation capacity of the resulting iPSC lines.

In this study, we set out to establish an optimized nonintegrative reprogramming approach under defined conditions in order to study the functional effects of transient p53 suppression in normal human dermal fibroblasts (NHDFs) during and after reprogramming.

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RESULTS

Increased Reprogramming Efficiency with Transient p53 Suppression

To study the effect of transient p53 suppression during reprogramming of human fibroblasts, a nonintegrative reprogramming system was established in defined conditions (Figure 1A and Figure S1 available online). Seven days after reprogramming with or without (w/wo) transient expression of a short hairpin to TP53 (shp53), a subpopulation of small, highly proliferative cells was observed in both conditions (Figures 1B and 1F). At days 14–28, iPSC colonies were observed (Figures 1C, 1D, 1G, and 1H), which stained positive for TRA-1-81 (Figures 1E and 1I). The growth rate was, in general, higher during reprogramming with transient p53 suppression (0.12 cell doublings per day) compared to without (0.06 cell doublings per day), although not to a significant degree (Figure 1J). In contrast, a fibroblast line, which stably expresses shp53 (PLK.O; Godar et al., 2008) displayed a significantly higher growth rate (0.42 cell doublings per day), yet, this line proved resistant to reprogramming (data not shown). Alkaline phosphatase staining and counting on day 28 showed a significant (7.5-fold) increase in the number of iPSC colonies with transient p53 suppression (0.12%) compared to without (0.016%) (Figure 1K), which was confirmed in several independent experiments comprising NHDFs from different individuals.

Transient p53 Suppression Increases Reprogramming Efficiency through Inhibition of p21 without Affecting Apoptosis and DNA Damage

To investigate the underlying effect of the increased reprogramming efficiency with transient p53 suppression, temporal changes in pluripotency markers (SSEA4, TRA-60), tumor suppression (p53), cell-cycle regulation (p21), apoptosis (PUMA), and DNA damage (H2A.X) were studied by flow cytometry. Although the PLK.O line stably expressing shp53 showed a 43.6% reduction in p53-positive cells compared to NHDFs, the pluripotency markers SSEA4 and TRA1-60 remained low during the entire reprogramming experiment (data not shown). In contrast, the analyses revealed a significant increase in SSEA4 and TRA1-60 double-positive cells with transient p53 suppression on day 21 (10.0% versus 3.1%) and on day 28 (16.6% versus 8.3%) after reprogramming (Figure 2A). A significant decrease in p53-positive cells was also observed with transient p53 suppression at days 7 (82.8% versus 90%), 14 (79% versus 92.7%), and 21 (63.8% versus 89.4%) (Figure 2B), which correlated with a significant decrease in p21-positive cells at days 7 (92.6% versus 96.3%), 14 (87.9% versus 95.6%), and 21 (68.1% versus 93.4%) (Figure 2C). In contrast, no significant effect of transient p53 suppression on apoptosis was observed, as evaluated by the proapoptotic marker PUMA (Figure 2D), a mitochondrial membrane assay (Figure S2A), and Annexin V in TRA1-60-positive cells (Figure S2B). Moreover, analysis with H2A.X, which is associated with replication-induced DNA damage (Garcia-Canton et al., 2012), detected no significant effect of transient p53 suppression on DNA damage, neither in the total cell population (Figure 2E) nor in TRA1-60-positive cells (Figures S2C and S2D).

Next, gene expression analysis of TP53 and its downstream DDR targets P21 and PUMA, as well as the pluripotency marker NANOG was performed. The silencing effect of shp53 resulted in a significant decrease in TP53 expression in unsorted cells 7 days after reprogramming (Figure S2E), and in P21 expression 14 days after reprogramming (Figure S2F), whereas expression of PUMA was unaffected (Figure S2G). In addition, no significant effect of shp53 suppression in TRA1-60-sorted cells was observed (Figures S2I–S2K). In contrast, the pluripotency marker NANOG was significantly upregulated in both unsorted and sorted cells 21 and 14 days after reprogramming with shp53, respectively (Figures S2H and S2L).

iPSCs Generated by Transient p53 Suppression Display Normal In Vitro Characteristics

To examine whether transient p53 suppression confers a negative effect on the long-term stability, six iPSC lines established w/wo transient p53 suppression were characterized in detail. Overall, there were no differences in morphology (Figures 3A and 3E). The pluripotency markers OCT4, NANOG, SSEA3, SSEA4, TRA1-60, and TRA1-81 were all detected by immunocytochemical staining (Figures 3B–3D and 3F–3H) and quantitative real-time PCR analyses showed a comparable hypomethylation of the pluripotency-associated genes OCT4, NANOG, SALL4, and ZFP42, DNMT3B, and TP53 and downregulation of P21 compared with NHDFs (Figure 3I). Moreover, integration analyses confirmed lack of expression from exogenous genes (Figure S3A). A comparable hypomethylation of the pluripotency-associated genes OCT4, NANOG, SALL4, and RAB25 and hypermethylation of the fibroblast-associated gene UBE1L was also observed (Figure 3J). Genome-wide transcriptome profiling showed no significant up- or downregulated genes in iPSC lines generated w/wo transient p53 suppression and analysis with PluriTest showed that all iPSC lines scored within the predefined pluripotency and novelty scores (Figure 3K). Finally, in vitro differentiation confirmed by the ability to generate embryoid bodies (Figures 3L, 3M, 3P, and 3Q), which stained positive for BetaIII-tubulin (TUJ1) (Figures 3N and 3R), smooth muscle actin (SMA), and alpha-fetoprotein (AFP) (Figures 3O and 3S).
Effects of Transient p53 Suppression on Human iPSCs

Figure 1. Increased Reprogramming with Transient p53 Suppression in Defined Conditions

(A) Timeline showing the reprogramming in defined conditions with the episomal plasmids hOSKUL or hOSKUL + shP53 with or without (w/wo) a short hairpin to p53 (shp53).

(B–D and F–H) Morphology of normal human dermal fibroblasts (NHDFs) and induced pluripotent stem cells (iPSCs) at days 7, 17, and 28 after reprogramming w/wo shp53.

(E and I) Tra-1-81 staining of primary iPSC colonies 28 days after reprogramming w/wo shp53.

(J) Growth rate of cells electroporated with hOSKUL, hOSKUL + shp53, or a fibroblast line stably expressing shp53 (PLK.0) at days 7, 21, and 28 after reprogramming.

(K) Alkaline phosphatase staining and counting of primary iPSC colonies at day 28 after reprogramming w/wo shp53.

Scale bars, 200 μm (B, C, F, and G) and 400 μm (D, E, H, and I). Y error bars depict SD of three independent experiments. *p < 0.05. See also Figure S1.
The genomic stability of fibroblasts and iPSC lines generated with and without transient p53 suppression was analyzed in detail by chromosome studies and copy number variation (CNV) analyses. All the lines presented normal karyotype and did not contain any microscopically visible structural or numerical abnormalities (46, XY) (Figure S3B). CNV analyses showed no significant differences between the numbers of CNVs in iPSC lines generated with and without transient p53 suppression, displaying an average of 1.66 genomic CNVs (four gains and six losses), which were not present in the donor NHDFs (Table S1). Three of the CNVs were detected in iPSC lines without transient p53 suppression, with one line containing two CNVs (K1), one line containing a single CNV (K3), and one line containing no CNVs (K2). In contrast, seven of the CNVs were detected in iPSC lines with transient p53 suppression, with one line (K1_shp53) containing five CNVs, whereas the other two lines (K2_shp53 and K3_shp53) had only a single CNV each, the latter in a noncoding region.

### DISCUSSION

In this study, we report the establishment of a nonintegrative reprogramming approach in defined conditions, based on transient suppression of p53. Using this method, we were able to consistently generate AP and TRA1-81-positive iPSC colonies from individuals with...
Stem Cell Reports

Effects of Transient p53 Suppression on Human iPSCs

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Effects of Transient p53 Suppression on Human iPSCs

different genders and ages. Compared to the results of Okita et al. (2011), who reported an average of 30 iPSC colonies per \( 1 \times 10^5 \) human fibroblasts (0.03%), we observed a 3-fold increase (0.11%) at defined conditions. In contrast, a fibroblast line stably expressing shp53 did not produce any iPSC colonies, and it is likely that the uncontrolled growth of this line has made it resistant to reprogramming due to accumulation of DNA damage or severe shortening of telomeres.

To study temporal changes during reprogramming with transient p53 suppression, flow cytometry with the pluripotency markers SSEA4/TRA1-60, which were previously used for isolation of fully reprogrammed iPSCs (Kahler et al., 2013), was performed. A significant increase in the amount of SSEA4/TRA1-60 double-positive cells was observed on day 21 (3-fold) and on day 28 (2-fold) after reprogramming with shp53. Moreover, transient p53 suppression also accelerated the temporal appearance of fully reprogrammed iPSCs, as the cells became earlier positive for SSEA4/TRA1-60.

As expected from the silencing effect of shp53, p53 was significantly suppressed during reprogramming with transient p53 suppression, which correlated with suppression of p21, whereas expression of the proapoptotic marker PUMA remained unchanged. These results imply that transient p53 suppression increases reprogramming by activating cell proliferation through p21 suppression, rather than by decreasing apoptosis (Figure 2F). Moreover, the effect of transient p53 suppression appears to occur in fibroblasts prior to reprogramming, because no significant differences in \( TP53, P21, \) and \( PUMA \) were observed in TRA1-60-sorted cells (Figures S2I–S2L). The latter is in agreement with a recent study, which demonstrated that embryonic stem cells (ESCs) possess a nonfunctional p53-p21 axis, in which p53 activates specific microRNAs, which inhibit p21 expression, thereby affecting cell-cycle regulation (Dolezalova et al., 2012). In contrast, NANOG was up-regulated with transient p53 suppression in both unsorted and TRA1-60-sorted cells; thus, NANOG could be a potential downstream target of p53. In addition, knockout of \( tp53 \) in mouse fibroblasts was recently shown to increase reprogramming through actions of p21 by promoting a mesenchymal-to-epithelial (MET) transition (Brosh et al., 2013), which could also apply in human.

Marion and colleagues previously reported that knockout of \( TP53 \) in human and mouse fibroblasts allows for efficient reprogramming at the expense of increased DNA damage, which was attributed to a decrease in apoptosis of DNA damaged cells (Marion et al., 2009). In contrast, we found that transient p53 suppression did not induce DNA damage or result in a decrease in apoptosis during reprogramming. The underlying mechanism is likely to constitute a combination of nonintegrative plasmids, which prevents excessive DNA damage, and a low background expression of \( TP53 \), which may be sufficient to sustain the apoptotic pathway.

To examine whether transient suppression of p53 had a negative effect on the long-term stability of iPSCs, we performed detailed characterization of iPSC lines established w/wo transient p53 suppression. All iPSC lines were highly similar with respect to expression of pluripotency markers and showed a comparable upregulation of \( TP53 \) and downregulation of \( P21 \) compared to NHDFs. Methylation analyses showed a comparable hypomethylation of the pluripotency-associated genes \( OCT4, NANOG, SALL4, \) and \( RAB25 \) and hypermethylation of the fibroblast-specific gene \( UBE1L \), as previously reported for iPSCs and ESCs (Nishino et al., 2011). Furthermore, genome-wide transcriptome analyses showed no significant up- or downregulated genes and analysis with PluriTest, which is a fast and ethically superior alternative to teratoma assays in mice (Buta et al., 2013), showed that all the iPSC lines clustered within the predefined pluripotency and novelty scores. Finally, in vitro
**Figure 4.** iPSCs Generated by Transient p53 Suppression Can Differentiate to Functional Neurons In Vitro

(A) Timeline showing the directed neural differentiation of induced pluripotent stem cell (iPSC) lines generated with or without (w/wo) a short hairpin to p53 (shP53).

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difficult to determine whether a given alteration is new, because the parental cells can also be analyzed (Laurent et al., 2011). The iPSC lines generated w/wo shp53 displayed normal karyotypes, and structural CNV analyses showed that they contained between zero and five CNVs (1.66 CNVs in average), which were not present in the parental NHDFs. These numbers are comparable with recent reports using single-nucleotide variants (SNVs) analyses that reported an average of five coding mutations, of which around half was preexisting variants (SNVs) analyses that reported an average of five coding mutations, of which around half was preexisting.

The present study demonstrates that transient p53 suppression increases reprogramming efficiency without affecting apoptosis and DNA damage. Furthermore, iPSC lines generated w/wo transient p53 suppression are identical with respect to their pluripotent phenotype, their mutational load, and their differentiation capacity, rendering the method suitable for in vitro studies on patient-specific disease pathology with a potential for clinical translation.

EXPERIMENTAL PROCEDURES

A complete description of experimental procedures can be found in Supplemental Experimental Procedures.

Reprogramming

Normal human dermal fibroblasts (NHDFs; Lonza) were electroporated with plasmids encoding hOCT4 or hOCT4 with a short hairpin to TP53 (shp53) in combination with hSOX2, hKLF4, hL-MYC, and hLIN28 (Addgene plasmids 27076, 27077, 27078 and 27080), abbreviated hOSKUL or hOSKUL + shp53, respectively (Okita et al., 2011, 2013) and cultured in mTeSR1 medium (STEM-CELL Technologies) and Matrigel-coated dishes (BD Biosciences) from day 7 to day 28.

Flow Cytometry and Apoptotic Measurements

Flow cytometry was performed with monoclonal antibodies against SSEA4 and TRA1-60 (BD Biosciences), p53, p21, H2A.X (Cell Signaling Technology), and PUMA (Novus Biologicals) on a FACSarray Bioanalyzer or a BD Accuri C6 (BD Biosciences). Unlabeled and isotype-labeled NHDFs were used as controls for gating.

Characterization of iPSC Lines

Six iPSC lines were established with hOSKUL (K1, K2, and K3) or hOSKUL + shp53 (K1_shp53, K2_shp53, and K3_shp53) and characterized at passage 10 according to established pluripotency criteria (Martí et al., 2013). Genome-wide transcriptome analysis was performed using the HT12 v4 BeadChip microarray (Illumina). DNA methylation analyses were performed with the Cells-to-CpG Bisulfite Conversion Kit (Life Technologies), and PCR-amplified sequences from ten clones per chromosomal region were examined. Primers are specified in Table S2.

Chromosome Studies and Copy Number Variation Analyses

Metaphase chromosomes were investigated with G-banding using standard procedures. Copy number variation (CNV) analysis was performed using the high resolution CytoScan HD chromosome microarray platform (Affymetrix), and data were processed using Affymetrix Chromosome Analysis Suite (ChAS) and manually corrected for false-positives hits.

(B–I) Phase contrast morphology of neuroepithelium (NE) at day 12 (B and F), neural progenitor cells (NPC) at day 21 (C and G), and neurons at day 35 (D, E, H, and I).

(J–Q) Immunocytochemistry of NPCs at day 21 with SOX2, NESTIN, and Hoechst (J and N) and VIMENTIN, OCT4, and Hoechst (K and O) and of neurons at day 35 with TUJI and Hoechst (L–P) and vGLUT and Hoechst (M and Q).

(R–U) Intracellular calcium kinetics in iPSC-derived neurons generated without (R and S) or with shp53 (T and U). Baseline fluorescence was recorded for 10 min before application of 300 μM glutamate/10 μM glycine, 25 mM K+, 100 μM GABA and 300 μM acetylcholine. The fluorescence was normalized to the first data point of each of the traces.

Scale bars, 200 μm (B–D, F–H, and J–Q) and 100 μm (E and I).
Stem Cell Reports
Effects of Transient p53 Suppression on Human iPSCs

Directed Neural Differentiation
Directed neural differentiation was carried out according to Shi et al. (2012). Primary antibodies are listed in Table S3.

Statistical Analyses
Statistical analyses comprised a two-tailed Student’s t test with *p < 0.05 considered significant.

ACCESSION NUMBERS
The GEO (http://www.ncbi.nlm.nih.gov/geo/) accession number for the microarray data reported in this paper is GSE48665. CNV raw data are available upon request.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, three figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2014.07.006.

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Mikkel A. Rasmussen, Bjørn Holst, Zeynep Tümer, Mads G. Johnsen, Shuling Zhou, Tina C. Stummann, Poul Hyttel, and Christian Clausen
Figures

**Figure S1.** Optimization of non-integrative reprogramming procedure, related to Figure 1. A) Survival and flow cytometry analyses of normal human dermal fibroblasts (NHDFs) one day after electroporation with a GFP plasmid using a Neon™ electroporation device at different conditions.

B-E) Phase contrast morphology and green fluorescent protein (GFP) fluorescence of NHDFs electroporated with a single pulse at 1600 V for 20 ms (Protocol #4; B, D) or two pulses at 1200 V for 20 ms (Protocol #14; C, E). Scale bars correspond to 200 µm.
Figure S2. Apoptosis, DNA damage and expression of P53, P21, PUMA and NANOG during reprogramming with or without (w/wo) a short hairpin to p53 (shp53), related to Figure 2. Normal human dermal fibroblasts (NHDFs) were analyzed on day 0, 7, 14, 21 (and 28) after reprogramming with the episomal plasmids hOCT4, hSOX2, hKLF4, hL-MYC and hLIN28 (hOSKUL) w/wo shp53.

A) Mitochondrial membrane potential assay with the dyes JC-1 and DAPI. In apoptotic cells, the mitochondrial potential collapses and the monomeric form of JC-1 shows green fluorescence. B) Flow cytometry of cells double positive for Annexin V and TRA1-60. C) Flow cytometry of cells double positive for the DNA damage marker H2A.X and TRA 1-60. D) Flow cytometry with H2A.X of NHDFs subjected to different doses of ultraviolet radiation. E-L) Quantitative real-time PCR analyses in the total cell population (E-H) or in TRA1-60 sorted cells (I-L) after reprogramming w/wo shP53. E, I) TP53, F, J) P21, G, K) PUMA and H, L) NANOG. Relative expression is shown as the fold change ($2^{-\Delta\Delta Ct}$) with NHDFs as reference and GAPDH as reference gene. Y error bars depict standard deviation of three independent experiments. *p<0.05.
Figure S3. Expression analysis of episomal vectors in iPSC lines and karyotyping of induced pluripotent stem cell lines, related to Figure 3. A) Reverse-transcriptase PCR at passage 10 with the endogenous and exogenous pluripotency genes LIN28, SOX2, and OCT4 in iPSC lines generated with or without a short hairpin to p53. Normal human dermal fibroblasts (NHDFs) transfected two days before with hOSKUL served as a positive control and a sample without template served as negative control. An iPSC line (SBI) generated by lentivirus was included as a reference. B) Representative karyotypes of the founder normal human dermal fibroblasts (NHDFs) and induced pluripotent stem cell (iPSC) lines generated with or without a short hairpin to p53 (shp53).
**Tables**

**Table S1.** Copy number variations in induced pluripotent stem cell lines, related to Figure 3.

| iPSC line | Type | Chr | Band | Start position   | End position   | Size (kb) | Genes within the CNV |
|-----------|------|-----|------|------------------|----------------|-----------|----------------------|
| K1        | Gain | 1   | q44  | 245500158        | 245787005     | 287       | KIF26B              |
| K1        | Gain | 3   | p12.2| 81531958         | 82762047      | 1230      | GBE1                |
| K3        | Loss | 2   | q33.3| 205758866        | 205984167     | 225       | PARD3B              |
| K1_shp53  | Gain | 5   | q34  | 167784954        | 167951642     | 167       | WWCI, RARS          |
| K1_shp53  | Loss | 7   | q11.22| 69672679        | 70023881      | 351       | AUTS2               |
| K1_shp53  | Loss | 7   | q31.1| 110709365        | 110842032     | 133       | IMMP2L, LRRN3       |
| K1_shp53  | Gain | 10  | q21.1| 53966718         | 54049359      | 83        | PRKG1               |
| K1_shp53  | Loss | 12  | q23.1| 100658499        | 100697244     | 39        | DEPDC4, SCYL2       |
| K2_shp53  | Loss | 13  | q31.3| 94337996         | 94506524      | 169       | GPC6                |
| K3_shp53  | Loss | X   | q21.33| 94049304        | 94129700      | 80        |                     |

1.Induced pluripotent stem cell line, 2. chromosome, 3. kilobases, 4. copy number variation, 5. database of genomic variance.

**Table S2.** Primers used for methylation analyses, related to Figure 3.

| Gene | Primer names | Primer sequences 5’-3’ | Product sizes (bp) | Products chromosomal extend |
|------|--------------|------------------------|--------------------|----------------------------|
| NANOG | ConNRR-F1 | ttATATTTtGATTTAAAAGTGGAAA | 298 | Chromosome 12 |
|       | ConNRR-R1 | aCAACCCTaCTCTAaCTCACaAaAAC | 701.877-702.174 | |
| POU5F1 | ConOCT4-3-F | AtTGTTTGGATGTAAAAGGt | 221 | Chromosome 6 |
| (OCT4) | ConOCT4-3-R | CCAaCTtaCTCTaCTTaaTaaCTCC | 31.080.784-31.080.564 | |
| RAB25 | ConRAB25-F1 | AGGGGtATTATTtTGTtATTAGTTG | 178 | Chromosome 1 |
|       | ConRAB25-R1 | TaaTaaCTCaCTCTCaCTCaTaaCTCC | 7.519.337-7.519.514 | |
| SALL4 | ConSALL4-F1 | GGttATATGTGTGtATGGtATGA | 110 | Chromosome 20 |
|       | ConSALL4-R2 | TCTTaattTaaaAAATTTACCCCT | 20.615.097-20.615.206 | |
| UBE1L | ConUBE1L-F1 | GtTTGGtTTGGTTTtTGGTTGtAt | 169 | Chromosome 3 |
Lower case 't' corresponds to a cytosine in the unconverted target sequence. Lower case 'a' corresponds to a guanine in the unconverted target sequence. *Genome Reference Consortium Human Build 37.

Table S3. Antibodies used for confirmation of pluripotency and in vitro differentiation, related to Figure 3 and 4.

| Antibody and host species         | Dilution | Company and catalog number       |
|-----------------------------------|----------|-----------------------------------|
| **Pluripotency**                  |          |                                   |
| rabbit anti-NANOG                 | 1:500    | Peprotech, 500-P236               |
| goat anti-OCT4                    | 1:500    | Santa Cruz, sc-8628               |
| rat anti-SSEA3                    | 1:100    | Biolegend, 330302                 |
| mouse anti-SSEA4                  | 1:100    | Biolegend, 330402                 |
| mouse anti-Tra-1-81               | 1:200    | Biolegend, 330702                 |
| mouse anti-Tra-1-60               | 1:200    | Biolegend, 330602                 |
| **In vitro differentiation**      |          |                                   |
| mouse anti-Smooth muscle actin (SMA) | 1:500 | DAKO, M0851                       |
| rabbit anti-Alpha-1-fetoprotein (AFP) | 1:500 | DAKO, A0008                       |
| mouse anti-Beta-III-tubulin (TUJI) | 1:4000  | Sigma-Aldrich, T8660              |
| mouse anti-SOX2                   | 1:100    | RD systems, MAB2018               |
| rabbit anti-NESTIN                | 1:4000   | Millipore, ABD69                  |
| mouse anti-VIMENTIN               | 1:500    | DAKO M0725                        |
| sheep anti-VGLUT1                 | 1:200    | Abcam, AB79774                    |
Supplemental Experimental Procedures

Unless otherwise stated, consumables and reagents were purchased from Sigma-Aldrich, St. Louis, MO, USA. Plasmids were purchased from Addgene, Cambridge, MA, USA.

Reprogramming of normal human dermal fibroblasts and establishment of iPSC lines

NHDFs (Lonza, CC-2511) from an 18-year-old male (XY NHDF) and a 32-year-old female (XX NHDF) were cultured in fibroblast medium consisting of Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine, penicillin and streptomycin (Pen/Strep) and 2 ng/ml basic fibroblast growth factor (bFGF; Miltenyi Biotech, Bergisch Gladbach, Germany).

At passage 5-6, the NHDFs from a 18-year old healthy male were trypsinized and $10^5$ cells were electroporated with a total of 1µg of the episomal plasmids hOCT4 (Addgene plasmid #27076) or hOCT4 with a short hairpin to TP53 (shp53; Addgene plasmid #27077) in combination with hSOX2 and hKLF4, (Addgene plasmid #27078) and hL-MYC and hLIN28 (Addgene plasmid #27080), abbreviated hOSKUL or hOSKUL + shp53, respectively. As a control, a fibroblast line from the same NHDF line was electroporated with a PLK.O-shp53 plasmid (Godar et al., 2008) and stable clones expressing shp53 were selected using puromycin. The electroporation was performed using a Neon™ electroporation device (Life Technologies, Carlsbad, CA, USA) with a single pulse at 1600 V for 20 ms or two pulses at 1200 V for 20 ms. Optimization of the electroporation procedure was performed with an episomal GFP plasmid (Addgene plasmid #27082).

Immediately after electroporation, the cells were transferred to a single well of a 6-well culture dish containing warm fibroblast medium without Pen/Strep, which was changed the day after to fibroblast medium with Pen/Strep. Seven days after electroporation, NHDFs were trypsinized and split 1:2 onto hESC-qualified Matrigel-coated dishes (BD Biosciences, Franklin
Lakes, NJ, USA) and cultured in mTeSR1 medium (Stem Cell Technologies, Vancouver, BC, Canada) in 5% O$_2$, 5% CO$_2$ in N$_2$ with the medium replenished every other day.

On day 28, six iPSC lines were established from the same XY NHDF donor using the plasmids $hOSKUL$ (named K1, K2 and K3) or $hOSKUL$ + shp53 (named K1_shp53, K2_shp53 and K3_shp53). The primary iPSC colonies were dissected out manually by needles and transferred to new Matrigel-coated 6-well dishes and cultured in mTeSR1. After a second manual passage, the iPSC lines were split 1:6 every 5-6 days with Dispase (Stem Cell Technologies, Vancouver, BC, Canada). At passage 10, the iPSC lines were harvested for subsequent analyses or frozen in CryoStem freezing medium (Stemgent, Cambridge, MA, USA).

**Alkaline Phosphatase and TRA1-81 live staining**

On day 28 after reprogramming, iPSC colonies were live stained for 30 min with a TRA1-81 antibody (Stemgent, Cambridge, MA, USA) or fixed in 4% paraformaldehyde (PFA) and stained for Alkaline phosphatase (AP) using an AP detection kit (Merck-Millipore, Billerica, MA, USA). The total numbers of AP-positive iPSC colonies were quantified using a stereomicroscope.

**Flow cytometry**

On day 0 (NHDFs) and on days 7, 14, 21 and 28 after reprogramming, cells were trypsinized and washed in flow buffer containing 1% bovine serum albumin (BSA) in Dulbecco’s Phosphate-Buffered Saline (DPBS). Cells were either incubated for 45 min with a combination of APC conjugated anti-SSEA4 (BD Biosciences, Franklin Lakes, NJ, USA) and PE conjugated anti-TRA-1-60 (BD Biosciences, Franklin Lakes, NJ, USA or fixed for 15 min with 4% PFA, incubated for 30 min in 90% ice-cold methanol and incubated with Alexaflour 488-conjugated anti-p53, anti-p21, anti-phospho-histone H2A.X (Ser139) antibody (Cell Signaling Technology, Danvers, MA, USA).
or DyLight 488-conjugated anti-PUMA (Novus Biologicals, Littleton, USA) and PE-conjugated anti-SSEA4 and 647-conjugated anti-TRA-1-60 (BD Biosciences, Franklin Lakes, NJ, USA). After washing twice in flow buffer, flow cytometric analysis was performed with a BD FACSarray Bioanalyzer or a BD Accuri C6 (BD Biosciences, Franklin Lakes, NJ, USA). Unlabeled and isotype labeled NHDF’s were used as controls for gating. As a positive control, NHDFs were exposed to different ultraviolet (UV) treatments at 254 nm, ranging from 0, 0.2, 0.5, 1 and 2 J/cm².

**Apoptotic measurements**

Apoptosis was measured by a mitochondrial membrane potential assay (Chemometec, Alleroed, Denmark). Briefly, cells were stained with the dyes JC-1 and DAPI. In healthy cells, JC-1 forms aggregates showing red fluorescence, whereas, in apoptotic cells the mitochondrial potential collapses and JC-1 localizes to the cytosol in its monomeric form showing green fluorescence. Quantification of green and red JC-1 fluorescence was carried out by use of a Nucleocounter NC-3000 (Chemometec Alleroed, Denmark).

**Immunofluorescence staining**

At passage 10, all iPSC lines were fixed with 4% PFA in PBS for 15 min and stained by standard immunofluorescence staining procedures. The primary antibodies (Supplementary table 1) were visualized with the secondary antibodies Alexa 488 or Alexa 594 diluted 1:400 (Life technologies, Carlsbad, CA, USA) and counterstained with Hoechst bisbenzimide 33258. Images were acquired on a Leica DMRB-fluorescence microscope (Leica Microsystems, Wetzlar, Germany).

**Quantitative real-time polymerase chain reaction (qRT-PCR)**

Total RNA was purified from unsorted and TRA1-60 sorted cells during reprogramming as well as
from six iPSC lines at passage 10 and NHDFs by RNeasy kit (Qiagen, Hilden, Germany).
Conversion to cDNA and quantitative real-time PCR (qRT-PCR) were performed using TaqMan RNA-to-C<sub>T</sub>™ 1-step kit (Life Technologies, Carlsbad, CA, USA). Primers for qRT-PCR (Life Technologies, Carlsbad, CA, USA) included SOX2, ZFP42, NANOG, POU5F1 (OCT4), DNMT3B, TP53, P21 (CDKN1A), PUMA (BBC3) and GAPDH. Relative quantification was calculated using 2<sup>ΔΔCT</sup> with NHDFs as a reference and GAPDH as reference gene. A commercial iPSC line (named SBI; System Biosciences, Mountain view, CA, USA), generated by viral transduction of human foreskin fibroblasts with OCT4, SOX2, KLF4 and C-MYC, was used as a positive control.

**In vitro differentiation**
Embryoid body (EB) formation was performed by transferring Dispase-treated clumps of iPSC to ultra-low attachment plates (Corning, Corning, NY, USA) in mTeSR1. After 2 days of culturing, the medium was changed to DMEM/F12 containing 20% knockout serum replacement (Life Technologies, Carlsbad, CA, USA), 1x non-essential amino acid, 2 mM L-glutamine, 0.1 mM 2-mercaptoethanol and 1% pen/strep. After 7 days, the EBs were plated on 0.1% gelatin-coated culture dishes and cultured in DMEM supplemented with 10% FBS, 2 mM L-glutamine and 1% pen/strep for up to three weeks. The cells were fixed for 15 min in 4% PFA for immunocytochemical analyses with antibodies of TUJI, SMA and AFP (Supplementary table 1).

**Karyotyping**
At passage 10, all iPSC lines and NHDFs were treated for 45 min with KaryoMAX colcemid (Life Technologies, Carlsbad, CA, USA) and harvested in fresh fixative containing 25% acetic acid and 75% methanol. Karyotyping was performed on G-banded metaphase chromosomes using standard
procedures. At least 10 metaphases were analyzed per sample with an approximate resolution of 550 to 600 bands per haploid genome.

**Genome wide transcriptome analysis**

For whole transcriptome microarray analysis, RNA was isolated from the iPSC lines K1 and K1_shp53 at passage 16, K2 and K2_shp53 at passage 12 and K3 and K3_shp53 at passage 13, as well as NHDFs and SBI iPSC with RNeasy Kit (Qiagen, Hilden, Germany). cDNA was hybridized to a HT12 v4 BeadChip microarray (Illumina, San Diego, CA, USA), which contains more than 47,000 probes for known human genes. Raw data were analyzed with the PluriTest algorithm (www.pluritest.org) or processed in Genome Studio data analysis software (Illumina, San Diego, CA, USA) using quantile normalization and a detection p-value of < 0.01 and subsequently analyzed in Multi Experiment Viewer (www.tm4.org) using a comparative t-test with Bonferroni correction and p < 0.05.

**Copy number variation (CNV) analyses**

DNA was purified from all iPSC lines at passage 10 and founder NHDFs using Wizard genomic DNA purification kit (Promega, Madison, WI, USA). CNV analysis was performed using the high resolution CytoScan HD chromosome microarray platform (Affymetrix, Santa Clara, CA, USA), which provides 750,000 polymorphic (SNP, single nucleotide polymorphism) and 1,900,000 non-polymorphic (CNV) markers. Raw data were processed using Affymetrix Chromosome Analysis Suite (ChAS) and manually corrected for false positive hits. To prevent false positive CNVs arising due to inherent microarray “noise”, gains and losses of > 30kb (and >30 consecutive probes) were taken into consideration (confidence limit > 90%). The identified CNVs were compared with the Database of Genomic Variants (DGV; http://projects.tcag.ca/variation/) and 1038 phenotypically
healthy samples run on the same platform to exclude common variants. The data were interpreted using the UCSC Genome Browser (http://genome.ucsc.edu). Data is available upon request.

**Directed neural differentiation**

Directed neural differentiation was carried out according to Shi and colleagues (Shi et al., 2012), with minor modifications. Briefly, iPSC lines generated w/wo shp53 (n=6) were cultured in mTeSR1 and ESC-qualified Matrigel until they reached 70-80% confluence. The medium was then changed to neural induction medium (NIM) consisting of DMEM/F12 and Neurobasal medium (1:1; Life Technologies, Carlsbad, CA, USA), 1 x N2 and 1 x B-27 minus Vitamin A (both from Life Technologies Carlsbad, CA, USA), 1mM Glutamax and Pen/Strep including the dual SMAD inhibitors LDN193189 (0.2 µM) and SB431542 (10 µM). After 10-12 days, the cells were split 1:3 with Dispase onto poly-L-ornithine (20ug/ml) and laminin (5 µg/ml)-coated dishes and cultured in neural maintenance medium (NMM) consisting of NIM without LDN193189 and SB431542. When rosette structures were observed, 20 ng/ml bFGF was added for 4 days followed by a second split (1:3) using Dispase. Hereafter, the cells were split 1:3 by Accutase upon confluence. At days 21 and 35, the cells were fixed in 4% PFA for immunocytochemical analyses with neuron-specific antibodies (Supplementary table 1) and on day 80, recordings of intracellular calcium kinetics were performed. Cells in 96-well plates were incubated for 1h at RT with the fluorescent calcium indicator, Calcium 5 (Molecular Devices, Sunnyvale, CA, USA) dissolved in Hanks Balanced Salt Solution (HBSS; Life Technologies). Following loading, recordings were performed with an FDSS 7000 fluorescence kinetics plate reader (Hamamatsu Photonics K.K., Hamamatsu city, Japan). The excitation wavelength was 480 nm; the emission wavelength was 540 nm and the sampling frequency 1Hz.