Pluripotent stem cells escape from senescence-associated DNA methylation changes

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Pluripotent stem cells evade replicative senescence, whereas other primary cells lose their proliferation and differentiation potential after a limited number of cell divisions, and this is accompanied by specific senescence-associated DNA methylation (SA-DNAm) changes. Here, we investigate SA-DNAm changes in mesenchymal stromal cells (MSC) upon long-term culture, irradiation-induced senescence, immortalization, and reprogramming into induced pluripotent stem cells (iPSC) using high-density HumanMethylation450 BeadChips. SA-DNAm changes are highly reproducible and they are enriched in intergenic and nonpromoter regions of developmental genes. Furthermore, SA-hypomethylation in particular appears to be associated with H3K9me3, H3K27me3, and Polycomb-group 2 target genes. We demonstrate that ionizing irradiation, although associated with a senescence phenotype, does not affect SA-DNAm. Furthermore, overexpression of the catalytic subunit of the human telomerase (TERT) or conditional immortalization with a doxycycline-inducible system (TERT and SV40-TAg) result in telomere extension, but do not prevent SA-DNAm. In contrast, we demonstrate that reprogramming into iPSC prevents almost the entire set of SA-DNAm changes. Our results indicate that long-term culture is associated with an epigenetically controlled process that stalls cells in a particular functional state, whereas irradiation-induced senescence and immortalization are not causally related to this process. Absence of SA-DNAm in pluripotent cells may play a central role for their escape from cellular senescence.

[Supplemental material is available for this article.]
senescence, (3) immortalization, and (4) reprogramming into induced pluripotent stem cells, which we refer to as iP-MSC. Subsequently, DNAm was analyzed using a novel Infinium HumanMethylation450 platform that can assay more than 480,000 CpG sites at single-base resolution (covering 99% of RefSeq genes and 96% of CpG islands) (Bibikova et al. 2011). We demonstrate that SA-DNAm changes are significantly enriched in developmental genes such as homeobox genes. Particularly, SA-hypomethylation occurs in intergenic regions or nonexpressed genes. Neither senescence induction by gamma irradiation, nor telomere extension influence SA-DNAm. In contrast, the majority of SA-DNAm is prevented by reprogramming into iP-MSC. Therefore, SA-DNAm changes do not resemble stochastic modifications that accumulate over time—they rather reflect a tightly regulated epigenetic program that is prevented by reprogramming into a pluripotent state.

Results

Long-term culture-associated DNA methylation changes

Mesenchymal stromal cells from human bone marrow were expanded until they entered growth arrest after 80 ± 25 d and 35.1 ± 5.7 cumulative population doublings (cPD) (Fig. 1A). At early passage, all cell preparations fulfilled the commonly used criteria for definition of MSC such as fibroblastoid growth, surface-marker expression (CD14−/CD29+, CD31−/CD34+, CD45−/CD73+, CD90+, and CD105+), and differentiation potential toward osteogenic, adipogenic, and chondrogenic lineages (Dominici et al. 2006). Senescent MSC acquired large and flat morphology and expressed senescence-associated beta-galactosidase (SA-β-gal). The immunophenotype was maintained throughout culture expansion, but MSC of late passages lost their adipogenic differentiation potential and revealed reduced osteogenic differentiation potential (Fig. 1B; Supplemental Fig. 1A–D). DNAm profiles were compared between MSC of early passage (P2 or P3; n = 5) and MSC of late passage just before entering senescence (P7–P16 according to inter-individual variation in long-term growth curves; n = 4). To focus on the most relevant SA-DNAm changes, we have only considered CpG sites with at least 20% differential methylation and a limma-adjusted P-value of <0.05. Under these stringent criteria 1702 CpG sites were SA-hypomethylated, whereas 2116 CpG sites were SA-hypomethylated (Fig. 1C; Supplemental Tables 1,2).

DNAm changes were not simultaneously acquired within genes, but most of them were restricted to circumscribed genomic locations. CpG sites were classified according to gene regions as described before (Sandoval et al. 2011; Supplemental Fig. 2). Unexpectedly, SA-CpGs were hardly located within 200 bp upstream of the transcription start site (TSS200; P < 10^-18; hypergeometric distribution) and in the first exon (P < 10^-7)—those regions where differential methylation has a major impact on gene expression. In contrast, SA-hypermethylation was particularly enriched in the gene body (P < 10^-7), 3'UTR (P < 10^-5), and intergenic regions (P < 10^-4),
whereas SA-hypomethylation was highly enriched in intergenic regions ($P < 10^{-5}$) (Fig. 1D). Analysis of SA-CpG sites in the context of CpG islands (CGI) revealed preferential occurrence of SA-hypermethylation in the 2-kb flanking regions of CGIs (shore region); $P < 10^{-5}$, whereas SA-hypomethylation hardly occurred in the context of CGIs at all ($P < 10^{-264}$) (Fig. 1E). SA-CpG sites did not exhibit clear association with subtelomeric or centromeric regions. We searched for differentially methylated regions (DMR) where 100 adjacent CpG sites on the BeadChip are regulated accordingly—only two genomic regions revealed such a highly significant enrichment of SA-DNAm changes ($P < 10^{-5}$): the HOX8 locus on chromosome 17 (SA-hypermethylated) and the keratin associated protein ($KRTAP$) locus on chromosome 21 (SA-hypomethylated; Supplemental Fig. 3). Overall, spatio-temporal DNAm changes were highly reproducible in independent samples.

Gene expression profiles from MSC of early passage (P3–P5; $n = 3$) and late passage (P10; $n = 3$) were compared by microarray technology. Signal intensity was used as an indicator for gene expression level and correlated with DNAm profiles. As expected, higher DNAm level within 200 nucleotides upstream of the transcription start site ($TSS200$) and first exon coincided with markedly reduced gene expression levels. However, the specific DNAm changes upon long-term culture were hardly reflected by differential gene expression (Supplemental Fig. 4) and similar results were observed when we matched SA-DNAm with our previously published gene expression data on long-term culture of MSC (Wagner et al. 2008). Overall, SA-hypomethylated CpG sites revealed a mean signal intensity that was significantly below the average of all genes (Fig. 1F). Thus, most of these genes seem to be neither expressed at early nor at late passages. Gene ontology analysis of genes with SA-hypermethylated CpG sites or SA-hypomethylated CpG sites revealed highly significant enrichment of genes in categories associated with cellular differentiation (Supplemental Table 3). Very similar results were observed when we focused on genes with SA-CpG sites within the promoter region ($TSS1500$, $TSS200$, and 5′UTR). Using gene set enrichment analysis we found the most significant overlap of either SA-hypermethylated or SA-hypomethylated CpG sites within genes harboring trimethylated lysine 9 of histone H3 ($H3K9me3$; $P = 2.6 \times 10^{-13}$ and $8.2 \times 10^{-13}$, respectively), and targets of Polycomb-group family members EED ($P = 2.1 \times 10^{-3}$ and $1.6 \times 10^{-7}$) and SUZ12 ($P = 9.3 \times 10^{-6}$ and $2.6 \times 10^{-7}$) in ChIP array data of human ESC (Lee et al. 2006; Ben-Porath et al. 2004). Furthermore, SA-hypomethylated CpG sites were also enriched in $H3K27$ targets of various cancer cells (Schlesinger et al. 2007), $P = 4.0 \times 10^{-6}$; (Kondo et al. 2008), $P = 2.4 \times 10^{-5}$; (Acevedo et al. 2008), $P = 5.1 \times 10^{-4}$. Enrichment of SA-hypermethylation or SA-hypomethylation was further analyzed with published Chip array data for $H3K9me3$ and $H3K27me3$ in human MSC from adipose tissue (Delbarre et al. 2010) and for the Polycomb repressive complex 2 (PRC2) targets in human MSC from bone marrow (Wei et al. 2011); these marks were highly significantly enriched in genes with SA-hypomethylated CpG sites (Fig. 1G). Taken together, particularly SA-hypomethylation preferentially occurred in regions with repressive $H3K9me3$ and $H3K27me3$ histone marks and in PRC2 targets—this may contribute to the very low expression of the corresponding genes.

Gamma-irradiation does not affect DNAm

Ionizing radiation (IR) of MSC induced very similar phenotypic changes as replicative senescence. Treatment with 15 Gy completely abrogated proliferation and the majority of cells stayed metabolically active for more than one month (Fig. 2A). After 7 d the cells acquired the typical senescent morphology. Furthermore, IR impaired differentiation toward adipogenic and osteogenic lineage as described above (Fig. 2B; Supplemental Fig. 5), transiently upregulated $CDKN1A$, $CDKN2A$, and $TP53$ (Supplemental Fig. 6) and other investigators demonstrated activation of the corresponding signaling cascades on a protein level (Cmielova et al. 2012). DNAm profiles were compared before and 7 d after treatment with 15 Gy ($n = 4$). None of the CpG sites revealed significant changes upon irradiation (limma-adjusted $P < 0.05$) (Fig. 2C). Accordingly, radiation-induced senescence does not seem to be associated with SA-DNAm changes. Subsequently, we measured telomere length by monochorme multiplex quantitative PCR (Cawthon 2009). As expected, telomere length decreased by long-term culture ($P = 0.01$), whereas irradiation did not affect telomere length. Despite phenotypic similarity of long-term culture-associated senescence and irradiation-induced senescence, the causative mechanism seems to be different.

Immortalization does not reverse SA-DNAm

Overexpression of $TERT$, the catalytic subunit of human telomerase, is known to extend the lifespan of primary cells, but it is generally insufficient for immortalization (Bodnar et al. 1998; Kiyono et al. 1998). We retrovirally transduced MSC with a $TERT$-IRES-GFP construct in comparison to GFP-control. Constitutive overexpression resulted in growth advantage of $TERT$-transduced cells (Fig. 3A), but this was only observed in one of five experiments, and the cells entered a senescent state after 5 additional cPD in comparison to GFP-control (Supplemental Fig. 7). Telomere length was moderately increased by TERT overexpression, whereas SA-DNAm changes were not affected (Fig. 3B,E).

Alternatively, we immortalized MSC with a lentiviral tetracycline-inducible expression system for TERT and SV40 large T antigen (SV40-TAg). In the presence of doxycycline (Dox) these cells could be passaged for more than 80 passages without any signs of senescence. However, removal of Dox from the culture medium resulted in immediate proliferation arrest, and within 10 d the cells acquired a typical senescent phenotype with SA-β-gal expression (Fig. 3C; Supplemental Fig. 8).

$q$RT-PCR analysis revealed that $TERT$ was

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**Figure 1.** Long-term culture-associated changes in MSC. (A) Long-term growth curves demonstrate that MSC reach a senescent state after 2–3 mo of culture expansion. (B) Cells at higher passage acquire morphological changes, senescence-associated beta-galactosidase staining, and loss of adipogenic differentiation potential (green: BODIPY-staining of lipid droplets; blue: DAPI) and decline in osteogenic differentiation potential (Alizarin red staining of calcium phosphate precipitates; size bars 100 μm). (C) Comparison of DNAm between early (P2–P3) and late (P7–P16) passages. CpG sites with relevant SA-hypermethylation (red: 1702 CpG sites) and SA-hypomethylation (green: 2116 CpG sites) are depicted (adjusted $P$-value < 0.05 and DNAm change >20%). (D) SA-DNAm changes were classified according to gene regions, and (E) in relation to CpG islands. (F) Signal intensity of gene expression microarrays was used as an indicator for gene expression level. In particular, genes with SA-hypomethylated CpG sites were significantly less-expressed than the average. (G) Genes with SA-DNAm changes were subsequently mapped to genes with $H3K9me3$, $H3K27me3$ (Delbarre et al. 2010), and PRC2 targets (Wei et al. 2011) in human MSC. Enrichment analysis revealed that particularly genes with SA-hypomethylated CpG sites have been associated with these repressive histone marks. As an alternative, we focused on the subset of promoter-associated CpG sites ($TSS1500$, $TSS200$, or 5′UTR; striped bars), and this analysis also revealed association of genes with SA-hypomethylated CpG sites with those genes that have repressive histone marks ($H3K9$, $P < 10^{-28}$; $H3K27$, $P < 10^{-11}$; $EZH2$, $P < 10^{-7}$).
Pluripotent cells escape SA-DNAm changes

After 3–4 wk colonies with typical ESC-like morphology emerged, expressing pluripotency markers such as POU5F1, NANOG, and POU5F1 (OCT4), SOX2, MYC, and KLF4 (Takahashi et al. 2007). MSC at passage 3 were retrovirally infected to express reprogramming factors was silenced and PluriTest analysis of those with a homeobox domain (Pfam, tatives for binding of POU2F1 (GFI1; P = 0.0025), growth factor independent 1 transcription repressor (GFI1; P = 1.4 × 10⁻⁴), and GATA2 (P = 3.5 × 10⁻⁴). Furthermore, they showed enrichment in targets of MIR323B (P = 2.1 × 10⁻⁵) and MIR10A/MIR10B (P = 4.6 × 10⁻⁴). In contrast, 1930 CpG sites were overlappingly hypomethylated at late passage (Fig. 4F), which were significantly enriched in genes with sequence-specific DNA-binding transcription factor activity (GO:3700; P < 10⁻¹²), particularly those with a homeobox domain (PIAM, P < 10⁻⁴). They were also enriched in genes with promoter regions, which comprise motives for binding of POUSF1 (P = 1.1 × 10⁻⁴), growth factor independent 1 transcription repressor (GFI1; P = 1.4 × 10⁻⁴), and GATA4 (P = 3.5 × 10⁻⁴). Furthermore, they showed enrichment in targets of MIR323B (P = 2.1 × 10⁻⁵) and MIR10A/MIR10B (P = 4.6 × 10⁻⁴). In contrast, 1930 CpG sites were overlappingly hypomethylated at late passage (Fig. 4F), and these sites were significantly enriched in G-protein-coupled receptors (GO:4930; P = 1.6 × 10⁻⁸) and in genes with binding sites for POUSF1, a class 6 homeobox protein (P = 0.0025), STAT5A (P = 0.016), and GATA2 (P = 0.017).

Reprogramming into iP-MSC prevents SA-DNAm changes

Subsequently, we analyzed SA-DNAm of induced pluripotent MSC (iP-MSC). MSC at passage 3 were retrovirally infected to express POUSF1 (OCT4), SOX2, MYC, and KLF4 (Takahashi et al. 2007). After 3–4 wk colonies with typical ESC-like morphology emerged, expressing pluripotency markers such as POUSF1, NANOgé, and TRA-1-80 (Fig. 4A), whereas the expression of retrovirally encoded reprogramming factors was silenced and PluriTest analysis of global gene expression profiles supported the notion that iP-MSC were pluripotent (Muller et al. 2011; Shao et al. 2012). Furthermore, embryoid bodies derived from iP-MSC comprised derivatives of all three embryonic germ layers. For comparison we utilized three widely used human ESC lines (H1, H9, and HES2). Telomere length differed markedly between these three ESC lines, whereas iP-MSC revealed a significantly higher methylation in iP-MSC (1842 of 2116 CpG sites), whereas 25% of the SA-hypermethylated CpG sites were significantly methylated in iP-MSC (2027 of 2116 CpG sites; P < 0.05). Conversely, 46% of the SA-hypermethylated CpG sites were significantly less methylated in iP-MSC in comparison to late passage MSC (796 of 1702 CpG sites) (Fig. 4D). Notably, the same tendency was also observed when we compared iP-MSC with MSC of early passage that have so far acquired relatively few SA-DNAm changes: 87% of the SA-hypomethylated CpG sites revealed a significantly higher methylation in iP-MSC (1842 of 2116 CpG sites), whereas 25% of the SA-hypermethylated CpG sites were significantly methylated (419 of 1702 CpG sites; Supplemental Fig. 11). Comparison of DNAm profiles of MSC and iP-MSC may be difficult because of the stark differences between the two cell types. Yet, it is remarkable that pluripotent stem cells hardly revealed SA-DNAm changes.

We reasoned that those SA-DNAm changes particularly might be relevant, which appear to be prevented by pluripotency. SA-CpG sites were filtered to be differentially methylated in MSC of late passage in comparison to (1) MSC of early passage, (2) iP-MSC, and (3) ESC (adjusted P-value < 0.05; DNAm change >20%). A total of 404 CpG sites were overlappingly hypermethylated in MSC of late passage (Fig. 4E), which were significantly enriched in genes with sequence-specific DNA-binding transcription factor activity (GO:3700; P < 10⁻¹²), particularly those with a homeobox domain (PIAM, P < 10⁻⁴). They were also enriched in genes with promoter regions, which comprise motives for binding of POUSF1 (P = 1.1 × 10⁻⁴), growth factor independent 1 transcription repressor (GFI1; P = 1.4 × 10⁻⁴), and GATA2 (P = 3.5 × 10⁻⁴). Furthermore, they showed enrichment in targets of MIR323B (P = 2.1 × 10⁻⁵) and MIR10A/MIR10B (P = 4.6 × 10⁻⁴). In contrast, 1930 CpG sites were overlappingly hypomethylated at late passage (Fig. 4F), and these sites were significantly enriched in G-protein-coupled receptors (GO:4930; P = 1.6 × 10⁻⁸) and in genes with binding sites for POUSF1, a class 6 homeobox protein (P = 0.0025), STAT5A (P = 0.016), and GATA2 (P = 0.017).
Figure 3. (Legend on next page)
When we used our epigenetic-senescence-signature, the predicted passage numbers for iP-MSC and ESC were close to noncultured cells, although they have been cultured for over many passages (Fig. 3D). Several other investigators have previously compared DNA changes upon reprogramming (Doi et al. 2009; Lister et al. 2009). Publicly available data sets with Infinium Human Methylation27 BeadChip were retrieved and tested with the Epigenetic-Senescence-Signature (Harnett et al. 2011; Nishino et al. 2011; Nazor et al. 2012)—these pluripotent samples were always estimated close to noncultured cells (Supplemental Fig. 12).

**Discussion**

SA-DNA changes are tightly coordinated and may entail functional consequences of cellular senescence. On the other hand, SA-DNA does not seem to be the immediate trigger of cellular senescence. Irradiation-induced senescence, although phenotypically very similar to replicative senescence, was not associated with SA-DNA changes. Furthermore, SA-DNA was not influenced by TERT overexpression or immortalization. We demonstrate for the first time that most of the SA-DNA changes do not occur in iP-MSC—and this supports the notion that these epigenetic modifications are functionally relevant. Our results indicate that long-term culture is associated with a process of cellular differentiation, which continuously locks specific genomic regions in an inactive state. Prevention of SA-DNA upon reprogramming into iPSC may be crucial for rejuvenation and escape from senescence (Fig. 5).

MSC resemble heterogeneous cell preparations—the composition of subpopulations is greatly affected by culture conditions and changes during culture expansion (Wagner and Ho 2007; Schellenberg et al. 2012). Therefore, future work should aim for analysis of SA-DNA changes on single cell level. On the other hand, similar SA-DNA changes were also observed in dermal fibroblasts (Koch et al. 2011, 2012), indicating that the observations made in this study are not restricted to MSC.

We describe how SA-CpG sites are enriched in the gene body, 3’UTR, and intergenic regions. CpG-methylation in these regions is not directly related to gene expression levels (Ball et al. 2009). Particularly, SA-hypomethylated CpG sites are associated with genes showing neither expression at early, nor at late passage. In fact, none of the genes of the Epigenetic-Senescence-Signature was detected by qRT-PCR (Koch et al. 2012). This might explain why overall the SA-DNA did not correlate with differential gene expression. Even though, SA-DNA changes may be functionally relevant to permanently lock genes in an inactive state. DNA changes result in tighter wrapping of DNA around the histone core (Lee and Lee 2012) and may support reorganization of the nuclear compartment by methyl-CpG-binding proteins (MBPs) (Sasai and Defossez 2009). It has been suggested that DNA does not have the primary determinant of gene activation in MSC but rather post-translational modifications of histones (Collas 2010; Sorensen et al. 2010). DNA may feedback on the histone code—potentially by changing the binding affinity of noncoding RNAs. PRC2 is recruited to target genes by noncoding RNAs such as HOTAIR, a transcript from the HOXC locus (Rinn et al. 2007); Xist, a non-coding RNA involved in X-chromosome inactivation (Zhao et al. 2008); and by the antisense-RNA 1 of CDKN2B (CDKN2B-AS1) (Yap et al. 2010). A recent comparison of epigenomic landscapes between human ESC and fibroblasts revealed that the repressive H3K9me3 and H3K27me3 marks expand upon differentiation into fibroblasts (Hawkins et al. 2010). In this study, we demonstrate that SA-hypomethylation, in particular, is significantly enriched in regions with H3K9me3, H3K27me3, and targets of PRC2, and is in line with previous observations (Schellenberg et al. 2011). It has been shown that aging induces DNA in PRC2 target genes and may thereby predispose to cancer (Teschendorff et al. 2010). Many cancers are derived from epithelial cells, and our previous work indicated that similar SA-DNA changes are also observed in other cell types including epithelial cells (Koch et al. 2012). This is of particular relevance, as SA-DNA changes might be involved in malignant transformation. In fact, hypomethylation of CpG sites that are highly methylated in ESC has recently been shown to be a key determinant of tumor invasion and clinical outcome (Zhuang et al. 2012).

The irreversible nature of senescent cells is generally believed to be due to alterations in the chromatin structure. Upon senescence, chromatin undergoes dramatic remodeling through the formation of facultative heterochromatin domains called senescence-associated heterochromatic foci (SAHF) (Narita et al. 2003, 2006). These compacted foci of DNA disappear only 7 d after reprogramming of senescent cells, and this was even considered as a first sign of iPSC generation (Lapasset et al. 2011). Further research will be necessary to demonstrate whether the SA-DNA reported here contributes to the formation of SAHF.

Ionizing radiation has been shown to induce a senescent phenotype in various different cell types (Di Leonardo et al. 1994; Igarashi et al. 2007); yet it is unclear whether the underlying molecular mechanisms are related to long-term culture-associated senescence (Rando and Chang 2012). We demonstrate that irradiation of MSC results in permanent proliferation arrest, typical senescent morphology, and loss of adipogenic and osteogenic differentiation potential. In contrast to replicative senescence, this stress-induced senescence was not associated with telomere attrition and did not evoke reproducible SA-DNA changes 7 d after irradiation. It is conceivable that epigenetic alterations are initially not present at the level of DNA, but rather on other levels of the epigenome: It has been suggested that factors involved in transcriptional repression and gene silencing are recruited to the...
site of DNA damage, including H3K9me3, H3K27me3, and repressive Polycomb complexes (O’Hagan et al. 2008; Chou et al. 2010; Seiler et al. 2011). This gene silencing may ultimately activate the same cell-intrinsic senescence program as described for cellular senescence—but the phenotypic and functional changes observed 7 d after irradiation are not causally related to DNA changes at specific sites in the genome.

Figure 4. Reprogramming into iP-MSC prevents SA-DNAm changes. (A) Induced pluripotent MSC reveal typical ESC-like morphology and express the pluripotency markers POU5F1, NANOG, and TRA-1-80. (B) Unsupervised hierarchical clustering of global DNA profiles clearly separates pluripotent cells (iP-MSC and ESC) and it discerns MSC of early and late passage. (C) Scatter plot of DNA profiles of iP-MSC in comparison to MSC of late passage (P7–P16). In particular, CpG sites that are hypomethylated upon replicative senescence (green) are hypermethylated in iP-MSC. In contrast, SA-hypomethylated CpG sites (red) are rather hypomethylated in iP-MSC. (D) Differential DNA methylation changes upon reprogramming into iP-MSC. Overall, SA-DNAm changes are prevented in iP-MSC. (E,F) Venn diagrams demonstrating the overlap of SA-DNAm changes with differential methylation between MSC of late passage with either iP-MSC or ESC.

Immortalization is often considered a counterpart of cellular senescence, but we demonstrate that it does not reverse SA-DNAm changes. As described by many other studies, overexpression of TERT may provide a proliferative advantage, but does not assure immortalization of MSC (Bodnar et al. 1998; Kiyono et al. 1998). In contrast, combination of TERT with SV40-TAg facilitates reliable immortalization (O’Hare et al. 2001). In this study, we describe for the first time a conditionally immortalized MSC line with TERT and SV40-TAg using a tetracycline-inducible system. Despite immortalization, the in vitro differentiation potential decayed upon long-term culture—adipogenic differentiation could not be induced at passage 13 and osteogenic differentiation decreased at higher passages. TERT overexpression, as well as TERT/SV40-TAg immortalization, resulted in extended telomeres, but they did not affect SA-DNAm. Therefore, immortalization of MSC does not address the regulatory mechanisms of SA-DNAm, and these cells may still undergo functional changes throughout culture expansion.

Telomere homeostasis is also crucial for pluripotent cells. Telomere shortening is a barrier for iPSC generation (Marion et al. 2009b), whereas transfection of TERT and SV40-TAg in combination with the classical four factors (POU5F1, SOX2, MYC, and KLF4) has been shown to enhance the reprogramming efficiency (Park et al. 2008). Telomere extension upon reprogramming may be due to telomerase expression, or it may be facilitated by alternative lengthening of telomeres (ALT), which is based on homologous recombination between telomeric sequences (Dunham et al. 2000; Lapasset et al. 2011). Epigenetic modifications such as H3K9me3, H4K20me3, and histone acetylation of telomeric chromatin have been shown to be involved in the regulation of telomere length (Benetti et al. 2007; Schoeftner and Blasco 2009). Furthermore, a decrease in DNA methylation is particularly at subtelomeric regions, is accompanied by dramatic elongation of telomeres (Gonzalo et al. 2006). SA-DNAm was not enriched in subtelomeric regions, and it is yet unclear whether or not it influences telomere length. To this end, our results demonstrate that telomere homeostasis is not the driving mechanism of SA-DNAm changes.

Replicative senescence has been described as a barrier to reprogramming of iPSC as the reprogramming efficiency is much higher in cells of early passage. A series of publications indicated that particularly up-regulation of TP53, CDKN1A, and CDKN2A may be responsible for this loss of reprogramming efficiency.
Expansion. Alternatively, SA-DNAm might even be reversed in the and that this low level is then maintained throughout culture with regard to cellular aging. It is therefore conceivable that our

nescence or whether it is a mere marker. MSC are heterogeneous—also proven whether SA-DNAm resembles the initial cause of senes-
culture stress (premature senescence). Furthermore, it is still not
demonstrated whether SA-DNAm changes are directly related to proliferation (replicative senescence) or rather some other type of
culture defects, but rather resembles an organized process that stalls cells in a particular functional state. It remains to be
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Figure 5. Schematic overview of how SA-DNAm changes may influence cellular senescence. Senes-
cence is associated with a continuous gain of SA-DNAm changes that are enriched in genes with re-
pressive histone marks. They may be associated with permanent gene silencing, but this link is yet
unproven and therefore labeled with a question mark. SA-DNAm changes are not affected by ionizing radiation or telomere length. However, these processes evoke chromosomal aberrations and may ul-
timately activate the same downstream signal cascades of cellular senescence (including activation of CDKN1A, CDKN2A, and TP53). Pluripotent cells hardly reveal SA-DNAm changes; avoidance from these epigenetic modifications might contribute to the escape of pluripotent cells from cellular senescence.

(Banito et al. 2009; Kawamura et al. 2009; Li et al. 2009; Marion et al. 2009a; Utikal et al. 2009). These factors are up-regulated upon senescence, and their ablation increased the reprogram-
ing efficiency. It is also conceivable that the donor age re-
sembles a barrier to reprogramming—other investigators dem-
strated that it is possible to generate iPSC from centenarians (Lapasset et al. 2011), and it will be of interest to systematically
analyze whether epigenetic rejuvenation is hampered in such samples. Overall, counteraction of senescence-associated modi-
fications seems to play a central role for the generation of iPSC. In
this regard, SA-DNAm might contribute to the reprogramming barrier at later passages.

Conclusion

All signs of cellular aging seem to be removed upon reprogram-
ing into iPSC. It has even been demonstrated that continuous passaging of iPSC further diminishes differences in DNA between iPSC and ESC (Nishino et al. 2011). Yet, upon rediffer-
entiation into fibroblasts they reveal typical signs of cellular aging, and finally enter a senescent state (Lapasset et al. 2011). Our
analysis reveals that SA-DNAm changes are tightly regulated. They are significantly enriched in genes with H3K9me3, H3K27me3, and PRC2-binding sites, indicating that SA-DNAm changes are
associated with heterochromatin formation. Overall, our results
support the notion that cellular aging is not only due to accumu-
lation of cellular defects, but rather resembles an organized process that stalls cells in a particular functional state. It remains to be
demonstrated whether SA-DNAm changes are directly related to proliferation (replicative senescence) or rather some other type of
culture stress (premature senescence). Furthermore, it is still not proven whether SA-DNAm resembles the initial cause of senes-
cence or whether it is a mere marker. MSC are heterogeneous—also with regard to cellular aging. It is therefore conceivable that our

In vitro differentiation

Adipogenic and osteogenic differentiation of MSC was induced as described before (Pittenger et al. 1999). After 18 d, osteogenic differentiation was analyzed by Alizarin Red staining. Adipo-
genic differentiated cells were stained with the green fluores-
cent dye BODIPY (4,4-difluoro-1,2,5,7,8-pentamethyl-4-bora-3a, 4a-diaza-s-indacene) and counterstained with DAPI (4’,6-Diamidin-2-phenylindol; both Molecular Probes). Quantification was performed as described in our previous work (Koch et al. 2011). Chondrogenic differentiation was induced in micromass culture for 18 d. Subsequently, the pellets were fixed with 10% formalin and first embedded in agarose and secondly in paraf-
fin; 1 mm sections were stained as previously described (Koch et al. 2011).

Senescence-associated beta-galactosidase

Expression of pH-dependent senescence-associated beta-galactosidase (SA-β-gal) was analyzed using the SA-β-gal staining kit (Cell Signaling Technology) or by flow cytometry with the fluorogenic substrate 5-dodecanoylaminofluorescein di-beta-D-galactopyranoside (C12FDG) (Debacq-Chainiaux et al. 2009).

Figure 5. Schematic overview of how SA-DNAm changes may influence cellular senescence. Senes-
cence is associated with a continuous gain of SA-DNAm changes that are enriched in genes with re-
pressive histone marks. They may be associated with permanent gene silencing, but this link is yet
unproven and therefore labeled with a question mark. SA-DNAm changes are not affected by ionizing radiation or telomere length. However, these processes evoke chromosomal aberrations and may ul-
timately activate the same downstream signal cascades of cellular senescence (including activation of CDKN1A, CDKN2A, and TP53). Pluripotent cells hardly reveal SA-DNAm changes; avoidance from these epigenetic modifications might contribute to the escape of pluripotent cells from cellular senescence.
Senescence induction by gamma-irradiation

MSC were treated with a blood product irradiator (model IBL-437, Institute for transfusion medicine, Aachen) with 5, 10, 15, 20, or 25 Gy as indicated in the text. Proliferation rate of MSC in early and late passage as well as after irradiation was determined in triplicate using the Alamar blue assay (Life Technologies), which was measured with a Tecan infinite M200 plate reader.

TERT overexpression

The TERT sequence was excised from the pGGRN145 plasmid (ATCC MBA-141) and subcloned into the retroviral expression vector pMIG (pMSCV-IRE-S-GFP) (Addgene plasmid 9044, Addgene) via EcoRI. MSC of five donors were transduced at passage 4 with the TERT-IRE-GFP construct, or for control with the corresponding empty pMIG vector (IRE-GFP), and the culture expanded in parallel until entering the senescent state.

Conditional immortalization with TERT/SV40-TAg

MSC were transfected with two lentiviral vectors (May et al. 2010): (1) SV40 large T antigen (SV40-TAg) under an unidirectional tetracycline-dependent promoter, which also drives a reverse transactivator through EMVC-IRE-S; (2) TERT under a bidirectional tetracycline-dependent promoter, additionally driving expression of an eGFP/neomycin phosphotransferase fusion gene linked by an EMVC-IRE element. Initially, MSC were transduced on 2 consecutive d with SV40-TAg and after 2 wk with TERT. Upon neomycin selection, the resulting colonies (~20 independent clones) were pooled and culture expanded in DMEM (PAA; 4.5 g/L glucose) supplemented with glutamine (PAA), penicillin/streptomycin (PAA), 15% FCS (PAA), and doxycycline hyclate (Dox; final conc. 2 μg/mL; Sigma, Taufkirchen). Proliferation rate with and without Dox was analyzed using the Alamar blue assay as described above.

Telomere length analysis

The average telomere length was estimated by monochrome multiplex Q-PCR as described before (Cawthon 2009; Ziegler et al. 2012). In brief, telomeres (T) and the beta-globin gene (S; single copy) were amplified on a MyIQ2 Two-color Realtime PCR Detection-System (Biorad). Analysis was performed in triplicate in parallel to serial dilutions of standard DNA. The relative telomere to single-copy gene (T/S) ratio for experimental DNA samples was calculated by dividing the copy number of the telomere template T by the copy number of the beta-globin template S. Amplification efficiency for the telomere and the beta-globin was between 90% and 100% in each run. Specificity of the amplification was confirmed by melt-curve analysis.

Generation and characterization of iP-MSC

MSC from three donors (passage 3) were infected on 2 consecutive d with equal amounts of pMXs-based retroviruses (Addgene carrying the POU5F1, Sox2, Klf4, and MYC genes (Takahashi et al. 2007). On day 3, the medium was changed to high-glucose DMEM (Gibco), 20% FCS, 1% nonessential amino acids (NAA), 1× penicillin/streptomycin, 1× L-glutamine, 0.1 mM β-mercaptoethanol (βME), and 50 ng/mL basic fibroblast growth factor (bFGF) (Peprotech) (Cai et al. 2010). On day 6 the cells were plated on gelatin-coated dishes containing irradiated CF1 murine embryonic fibroblasts (MEFs), and on day 7 the culture medium was supplemented with 20 μg/mL vitamin C (Sigma) and 1 mM valproic acid (Sigma). Between days 21 and 30 the ESC-like colonies were observed and mechanically isolated. Established iP-MSC were maintained on MEFS in DMEM/F12 medium supplemented with Glutamax, 20% knockout serum replacer, 1% NAA, 0.1 mM βME, and 50 ng/mL bFGF. Cells were passaged by manual dissection of cell clusters every 5–6 d. Human ESC lines H1, H9, and HES-2 (WiCell Research Institute) were used as controls and cultured under the same conditions as described above. Work with human ESC has been approved by the Robert Koch Institute (permit no. Az 1710-79-1-4-57). Immunohistochemical analysis of pluripotency markers was performed with primary antibodies against POU5F1 (Santa Cruz), SOX2 (Stemgent), NANOG (Stemgent), or TRA-1-80 (Santa Cruz) by alkaline phosphatase staining (Sigma) and by flow cytometric analysis of TRA-1-60 (BD Pharomingen) and SSEA4 (Santa Cruz). Furthermore, iP-MSC differentiation was analyzed using the embryoid body differentiation system with immunostaining after 14–19 d for desmin (mesoderm), alpha-fetoprotein (endoderm), and nestin (ectoderm). Generation of iP-MSC clones and analysis of pluripotency has been described in further detail elsewhere (Shao et al. 2012).

DNA methylation analysis

Genomic DNA was isolated from 10⁶ cells using the Qiagen DNA Blood Midi-Kit. Bisulfité-converted DNA (EZ DNA Methylation Kit; Zymo) was prepared and then analyzed using the InfiniumHumanMethylation450 BeadChip (Illumina) according to the manufacturer’s instructions. DNAm is provided as beta-values for each of the 485.577 CpG sites, ranging from 0 (nonmethylated) to 1 (100% methylation). Hybridization and initial data analysis with the BeadStudio Methylation Module was performed at the DKFZ Gene Core Facility in Heidelberg. Distribution of samples across the BeadChips is presented in Supplementary Table 4. Data have been deposited at NCBI’s Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo; accession no: GSE37066).

In addition, we analyzed all samples using our Epigenetic-Senescence-Signature, which has been described in detail before (Koch et al. 2012). This method is based on continuous DNAm changes at six specific CpG sites (associated with the genes GRM7, CASR, FRAMEF2, SELF, CASP14, and KRTAP13-3). DNAm at these specific CpG sites was analyzed by pyrosequencing (Varionostic GmbH) and subsequently used for linear regression models to estimate passage numbers, cPD, and days in culture. These calculations can also be performed with the online calculator, which is accessible under http://www.molcell.rwth-aachen.de/dms/.

Bioinformatic analysis of DNAm profiles

For further analysis we have only considered CpG sites on autosomes. Initially, we considered various normalization regimens, but we then decided to use raw data beta-values due to the global changes of DNAm upon reprogramming with an overall increase of DNAm level. Histograms of DNA methylation levels and principal components analysis (PCA) were calculated with R. Unsupervised hierarchical clustering according to Euclidian Distance (with Ward’s Minimum Variance Method) was performed using the MultiExperiment Viewer (MeV, TM4.7.4; http://www.tm4.org/mev/) or R. A DNAm changes were calculated by the limma package in R with adjusted P-values <0.05 and >20% change in DNAm level between early and late passages. Affiliation of CpG sites with gene regions or CGI was used as described in detail previously (Sandoval et al. 2011). Gene Ontology analysis of genes with DNAm changes was performed using the GoMiner software (http://discover.nci.nih.gov/gominer/) and enrichment of chromosomal location was analyzed with the gene set enrichment analysis (GSEA; http://
www.broadinstitute.org/gsea/index.jsp). Furthermore, we compared DNAm profiles with published chromatin immunoprecipitation (ChIP) results: data for H3K9me3 and H3K27me3 in adipose tissue-derived MSC were retrieved from GSE17053 (Delbarre et al. 2010) and further processed by NimbleScan; EZH2 targets in bone marrow MSC were retrieved from the supplemental material of another manuscript (Wei et al. 2011). For comparison, DNAm and ChIP data were matched by GenBank Accession IDs and enrichment of SA-DNAm in these regions was determined by hyper-geometric distribution.

Gene expression analysis

For gene expression analysis, three additional MSC samples were isolated and expanded as described above. Re-seeding was performed at a density of 1.3333 cells/cm² in standard medium consisting of DMEM (PAA; 1 g/L glucose) supplemented with glutamine (PAA), penicillin/streptomycin (PAA), and 10% FSC (PAA). RNA was isolated at early passage (P3, P4, or P5) and late passage (P10) with the miRNeasy kit (Qiagen). RNA concentration and integrity were determined by an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc.). Gene expression analysis was performed on GeneChip Human Gene 1.0 ST Arrays (Affymetrix) at the Interdisciplinary Center for Clinical Research (IZKF) at the RWTH Aachen, and subsequently projected to Plurifit analysis as described below (Muller et al. 2011). All samples were clearly associated with pluripotent cells and data are accessible at GEO Series GSE38806.

Data access

The microarray data from this study have been submitted to the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo) under accession numbers GSE37065, GSE37066, and GSE38806.

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