Senescence impairs successful reprogramming to pluripotent stem cells

Ana Banito,1 Sheikh T. Rashid,2 Juan Carlos Acosta,1 SiDe Li,3,4 Carlos F. Pereira,3 Imbisaat Geti,2 Sandra Pinho,6 Jose C. Silva,7 Veronique Auzuara,6 Martin Walsh,3,4 Ludovic Vallier,2 and Jesús Gil1,8

1Cell Proliferation Group, MRC Clinical Sciences Centre, Faculty of Medicine, Imperial College, London W12 0NN, United Kingdom; 2Laboratory for Regenerative Medicine, University of Cambridge, Department of Surgery, Cambridge CB2 0SZ, United Kingdom; 3Department of Pediatrics, Mount Sinai School of Medicine, New York, New York 10029, USA; 4Department of Structural and Chemical Biology, Mount Sinai School of Medicine, New York, New York 10029, USA; 5Lymphocyte Development Group, MRC Clinical Sciences Centre, Faculty of Medicine, Imperial College, London W12 0NN, United Kingdom; 6Institute of Reproductive and Developmental Biology, Faculty of Medicine, Imperial College, London W12 0NN, United Kingdom; 7Wellcome Trust Centre for Stem Cell Research, University of Cambridge, Cambridge CB2 1QR, United Kingdom; 8Corresponding author.

Somatic cells can be reprogrammed into induced pluripotent stem (iPS) cells by overexpressing combinations of factors such as Oct4, Sox2, Klf4, and c-Myc. Reprogramming is slow and stochastic, suggesting the existence of barriers limiting its efficiency. Here we identify senescence as one such barrier. Expression of the four reprogramming factors triggers senescence by up-regulating p53, p16INK4a, and p21CIP1. Induction of DNA damage response and chromatin remodeling of the INK4a/ARF locus are two of the mechanisms behind senescence induction. Crucially, ablation of senescence effectors improves the efficiency of reprograming, suggesting novel strategies for maximizing the generation of iPS cells.

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The ability to derive pluripotent cells from somatic cells may fulfill the promise of unlimited stem cells for applications in basic biology, drug development, or regenerative medicine. Different strategies have been used to reprogram somatic cells into pluripotent stem cells. These include nuclear transfer [Wilmut et al. 1997], cellular fusion with embryonic stem (ES) cells [Ying et al. 2002], or the expression of factors linked to stem cell pluripotency [Takahashi and Yamanaka 2006]. Regarding the latter, Takahashi and Yamanaka [2006] first showed that mouse somatic cells could be reprogrammed to a pluripotent-like state by expressing four transcription factors (Oct4, Sox2, Klf4, and c-Myc). Since then, induced pluripotent stem (iPS) cells have been derived from cells of multiple origin, including human adult fibroblast and keratinocytes [Takahashi et al. 2007; Yu et al. 2007; Lowry et al. 2008; Park et al. 2008]. Alternating the combination of factors [i.e., substituting Klf4 and c-Myc by Nanog and Lin28] has been used successfully to reprogram different cells or improve the efficiency of reprogramming [Yu et al. 2007; Feng et al. 2009]. Currently, the two major limitations for translating iPS-based reprogramming into clinical application are to routinely deliver iPS cells safe and free of vector and transgene sequences, and to improve the reprogramming efficiency. The first issue is being addressed by using alternative methods to deliver the reprogramming factors such as transient transfection [Okita et al. 2008], transposons [Kaji et al. 2009], adenoviral [Stadtfeld et al. 2008], episomal vectors [Yu et al. 2009], or recombinant proteins [Zhou et al. 2009]. Different strategies are under investigation to enhance the efficiency of reprogramming [Feng et al. 2009].

Reprogramming occurs as a gradual process; for example, upon retroviral transduction, lineage-specific genes are silenced gradually and, as embryonic markers become activated, endogenous genes linked to pluripotency are induced [Brambrink et al. 2008]. However, this process results in only a small percentage of the infected population of cells becoming pluripotent. This suggests that barriers could be limiting the efficiency of successful reprogramming. The comparison of genetic and epigenetic profiles of partially reprogrammed cells (Pre-iPS) and iPS show that independently derived Pre-iPS share similar gene expression and epigenetic profiles, reflecting a common intermediate state [Mikkelsen et al. 2008; Sridharan et al. 2009]. Pre-iPS display incomplete epigenetic remodeling and persistent DNA hypermethylation, and can be converted into iPS with varied efficiency through global inhibition of DNA methylation [Mikkelsen et al. 2008]. The existence of other mechanisms limiting reprogramming efficiency remains to be investigated.

Senescence is the irreversible arrest during the G1 transition of the cell cycle elicited by replicative exhaustion or in response to stresses such as DNA damage, chemotherapeutic drugs, or aberrant expression of onco-genes. This arrest is implemented primarily through activation of p53 and the up-regulation of the cyclin-dependent kinase (CDK) inhibitors p16INK4a and p21CIP1 [Collado et al. 2007]. Beyond being simply an in vitro observation, senescence plays relevant physiological roles. In particular, oncogene-induced senescence (OIS) is widely observed among varied premalignant lesions and is believed to be tumor-suppressive. Senescence also limits the homeostasis of normal tissues through the regulation of stem cell self-renewal, therefore impacting on aging [Collado et al. 2007].

In this report, we investigate the direct relationship between the process of iPS reprogramming and senescence. We observe that reprogramming initially triggers a stress response with characteristics of senescence. This reprogramming-induced senescence (RIS) acts as an
Reprogramming-induced senescence

Results and Discussion

We analyzed the early response of somatic cells to expression of the four reprogramming factors (Oct4, Sox2, Klf4, and c-Myc) [Fig. 1A]. To this end, a polycistronic cassette encoding Oct4, Sox2, Klf4, and c-Myc (OSKM) [Carey et al. 2009] was transferred to a retroviral vector [pBABE-OSKM]. Early events after the infection of IMR90 human fibroblasts with the OSKM vector [Fig. 1] or coinfection with four individual vectors expressing the reprogramming factors [4-F] [Supplemental Fig. S1] were monitored. As controls, we included cells infected with an activated version of RAS [H-RASG12V], able to cause senescence. Upon expression of the reprogramming factors, we observed a decrease in the numbers and growth of IMR90 cells as evaluated by growth curves, colony formation assays, and a decrease in the percentage of cells incorporating BrdU [Fig. 1B–D]. Expression of the reprogramming factors increased the percentage of cells arrested in G1, without inducing apoptosis significantly in our experimental conditions [Supplemental Fig. 1]. Cells expressing the four reprogramming factors resembled senescent cells with an enlarged cytoplasm and displayed senescent-associated β-galactosidase (SA β-Gal) activity, and senescence-associated heterochromatic foci (SAHF) [Fig. 1E; Supplemental Fig. 1].

Expression of reprogramming factors in another strain of human fibroblasts (BJ) or in mouse embryonic fibroblasts (MEFs) (Supplemental Fig. S5A) was transferred to a retroviral vector (pBABE-OSKM). Early events after the infection of IMR90 human fibroblasts with the OSKM vector [Fig. 1] or coinfection with four individual vectors expressing the reprogramming factors [4-F] [Supplemental Fig. S1] were monitored. As controls, we included cells infected with an activated version of RAS [H-RASG12V], able to cause senescence. Upon expression of the reprogramming factors, we observed a decrease in the numbers and growth of IMR90 cells as evaluated by growth curves, colony formation assays, and a decrease in the percentage of cells incorporating BrdU [Fig. 1B–D]. Expression of the reprogramming factors increased the percentage of cells arrested in G1, without inducing apoptosis significantly in our experimental conditions [Supplemental Fig. 1]. Cells expressing the four reprogramming factors resembled senescent cells with an enlarged cytoplasm and displayed senescent-associated β-galactosidase (SA β-Gal) activity, and senescence-associated heterochromatic foci (SAHF) [Fig. 1E; Supplemental Fig. 1]. Expression of reprogramming factors in another strain of human fibroblasts (BJ) or in mouse embryonic fibroblasts (MEFs) [Supplemental Fig. S5A] also resulted in a growth arrest with characteristics of senescence [Supplemental Fig. 2]. To dissect how each of the reprogramming factors contributed to senescence, we infected IMR90 cells with each factor separately. To our surprise, each of the four factors—Oct4, Sox2, Klf4, and cMyc—decreased cell growth, decreased BrdU incorporation, and up-regulated markers and characteristics of senescence in IMR90 cells [Supplemental Fig. S3], thus suggesting a complex scenario in which different pathways could be activated simultaneously to trigger senescence in response to their combined expression.

To identify the pathways responsible for senescence induction at the early stages of reprogramming, IMR90 cells were infected with the OSKM vector and analyzed by immunofluorescence. Expression of reprogramming factors caused a DNA damage response as observed by elevated levels of nuclei staining positive with the pST/Q or γH2AX antibodies [Fig. 2A; data not shown]. Interestingly, OSKM expression also caused oxidative stress, as reflected by elevated levels of the oxidized base 8oxoG [Supplemental Fig. S5A]. We also observed a significant up-regulation of the tumor suppressors p16INK4a, p53, and p21Cip1—critical senescent effectors—upon infection with the OSKM polycistronic vector in IMR90 cells [Fig. 2B–D; Supplemental Fig. 5], BFs, and MEFs [Supplemental Fig. S2]. Reanalysis of published transcriptional profiles of MEFs expressing the four factors under inducible control [Mikkelsen et al. 2008] showed increased p16INK4a and p21Cip1 levels that decline slightly at later times [Supplemental Fig. S6]. A time course in IMR90 cells expressing the reprogramming factors showed that although levels of senescence effectors are elevated [compare with vector], at later times their expression decreases, especially for p53 and p21Cip1 [Supplemental Fig. S7].

Interestingly, we also observed the up-regulation of p16INK4a and p21Cip1 [Supplemental Fig. S8C] during heterokaryon-based reprogramming [Pereira et al. 2008], suggesting an inherent link between senescence and reprogramming. We reasoned that if up-regulation of senescence effectors was an early response to reprogramming, their levels could remain elevated in Pre-iPS cells. Indeed, expression of p16INK4a and p21Cip1 was up-regulated in Pre-iPS derived from MEFs when compared with controls [Supplemental Fig. S9A], and similar conclusions were drawn from re-examination of published data [Supplemental Fig. 9B,C; Mikkelsen et al. 2008; Sridharan et al. 2009]. Therefore, induction of senescence upon reprogramming [RIS] is observed in different cell types, using different reprogramming approaches, and it is reflected by elevated levels of senescence effectors such as p21Cip1 and p16INK4a in Pre-iPSs.

To analyze the contribution of the senescence effectors up-regulated during RIS, we coexpressed the OSKM vector with the E6 and E7 proteins of HPV16, which inactivate the p53 and the Rb networks, respectively. As shown in Figure 2E, expression of either alleviates the growth arrest induced by the reprogramming factors, and joint expression of both E6 and E7 at least had additive effects. To test specifically for the contribution of initial barrier limiting the efficiency of the process. De- cisively, alleviating senescence in these cultures enhances reprogramming efficiency.

Figure 1. Induction of senescence upon expression of the four reprogramming factors. (A) Timeline of the experiments in this figure and Figure 2. IMR90 human fibroblasts were infected with a polycistronic vector expressing the four factors (Oct4, Sox2, Klf4, and c-Myc). Cells were selected for 7 d. (B) Effect of the expression of the four factors on the growth of IMR90 cells. (C) Crystal violet-stained plates of IMR90 cells infected with the indicated vectors. (D) Expression of the four reprogramming factors causes a decrease in the percentage of IMR90 cells incorporating BrdU. (E) Effect of the vector expressing the four reprogramming factors over SA β-Gal activity and SAHF formation in IMR90 cells.
p16INK4a, p21CIP1, and p53 to the arrest, we used shRNAs. Analysis of the BrdU incorporation and growth in cells infected with the OSKM vector showed that p53, p21CIP1, and p16INK4a are relevant for the arrest (Fig. 2E,F).

The INK4a/ARF locus is normally subjected to strong epigenetic repression mediated by H3K27 methylation and recruitment of Polycomb-repressive complexes. Therefore, we investigated the extent of H3K27 methylation at the INK4a/ARF promoter in response to the expression of reprogramming factors. Chromatin immunoprecipitation (ChIP) was used to measure the ratio of trimethylated H3K27 (H3K27me3) to total H3 using primer sets described previously (Fig. 3A; Barradas et al. 2009). The repressive mark peaked around the INK4a promoter, and we observed a decrease in the levels of H3K27me3 histone methylase EZH2 to the locus but only marginal effects on their total levels of expression (Fig. 3F; data not shown). We observed similar remodeling of the INK4a/ARF locus when the reprogramming factors were expressed from individual vectors (4-F) (Supplemental Fig. S10).

To understand how the p53/p21CIP1 pathway is engaged during RIS, we expressed each of the reprogramming factors individually. Sox2, Klf4, and c-Myc each up-regulated p21CIP1 levels (Fig. 4A). Interestingly, they induced p21CIP1 by different mechanisms. Sox2 expression resulted in a p53-independent up-regulation of p21CIP1, while c-Myc or Klf4 induced both p53 and p21CIP1, although only c-Myc provoked DNA damage.
negative cell cycle regulators such as p21CIP1, the Rb homolog p130, or LATS2 (Wang et al. 2008). In this regard, we noted that not only p21CIP1 but also the expression of p130 increased during RIS (Supplemental Fig. S11A). Several microRNAs [miRNAs] have been identified whose expression is linked to pluripotency, such as the miR-290 cluster in mouse cells, the miR-371–373 cluster in human cells, or the miR-302 cluster conserved in both [Houbaviy et al. 2003; Suh et al. 2004]. These miRNAs are expressed in ES cells and induced during the reprogramming of zygotes to the eight-cell embryo stage (Tang et al. 2007). Some of these miRNAs are necessary for the normal proliferation of ES cells, and function by targeting negative cell cycle regulators such as p21CIP1, the Rb homolog p130, or LAT52 [Wang et al. 2008]. In this regard, we noted that not only p21CIP1 but also the expression of p130 increased during RIS [Supplementary Fig. S11A].

The expression of the four reprogramming factors in IMR90 cells did not reset the level of these miRNAs to the ones observed in hES cells [Fig. 4B, Supplementary Fig. S12], even if the miR-302 cluster is regulated by Oct4 and Sox2 [Barroso-del Jesus et al. 2009]. In contrast, iPS cells expressed these miRNAs at levels comparable with those of hES cells [Fig. 4B, Supplementary Fig. 12]. Uncoupling between the expression of the four reprogramming factors and pluripotency-associated miRNAs may account for elevated p21CIP1 expression during RIS. In accordance with our hypothesis, ectopic expression of the miR-302 cluster alleviated the growth arrest [Fig. 4C] and prevented the up-regulation of p21CIP1 expression during RIS [Fig. 4D, E]. A corollary of our observations is that inhibition of senescence could improve reprogramming efficiency. To support this contention, we knocked down the expression of p16Ink4a, p19Arf, p21CIP1, or p53 in BJ human fibroblasts with shRNAs [Fig. 5A]. Next, we infected these cells with viruses expressing Oct-4, Sox2, Klf-4, and c-Myc and cultured them in the appropriate conditions to promote the appearance of human iPS [hIPs] cell colonies. After 21 d, colonies were analyzed for expression of NANOG and TRA-1-60 by immunofluorescence [Supplementary Fig. 13A]. Double-positive colonies were counted as fully reprogrammed iPS cell colonies, whereas morphologically distinct colonies negative for both markers were scored as partially reprogrammed [Fig. 5B, Supplementary Fig. S13A–C]. Similar experiments were performed with IMR90 cells using morphological criteria to determine hiPS cell colonies and partially reprogrammed colonies. Knockdown of senescence effectors resulted in an increased number of fully and partially reprogrammed iPS cell colonies in both BJ and IMR-90 cells [Fig. 5B, Supplementary Fig. S13]. Experiments carried out in MEFs in which the expression of p16Ink4a/p19Arf, p21CIP1, or p53 was knocked down by shRNAs, or in strains knocked out for p53 or p21, also revealed increased reprogramming efficiency upon depletion of senescence effectors [Supplementary Fig. S14].

hiPS cell colonies generated from IMR90 or BJ fibroblasts with reduced levels of senescent factors were individually picked and expanded. All the hiPS cell lines [n = 8] analyzed to date expressed pluripotency markers including Oct4, Sox2, Nanog, and Tra-1-60 [Fig. 5C,D; Supplementary Fig. S15], and the transgenes used for reprogramming were silenced in iPS [Fig. 5D]. The hiPS lines analyzed to date [n = 8] were capable of differentiating in vitro into extraembryonic tissues and into derivatives of the three germ layers—ectoderm, mesoderm, and endoderm [Fig. 5E]. Taken together, these data demonstrate that inhibition of senescence improves the efficiency of reprogramming of somatic cells, and the resulting iPS cells display characteristics of pluripotent stem cells.

Different reports have characterized cells that failed to fully reprogram [Pre-iPS] and suggest that they are trapped in a late step of reprogramming [Mikkelsen et al. 2008]. Inhibition of DNA methylation, knockdown of lineage-specific genes, or treatment with two inhibitors [Silva et al. 2008] can either convert some of these Pre-iPS to iPS or increase the proportion of fully reprogrammed iPS versus Pre-iPS. By inhibiting or alleviating senescence, we presumably increase the cells that surpass the early barrier imposed by RIS, resulting in higher numbers
of both Pre-iPS and fully reprogrammed iPS (Supplemental Fig. S16). A combination of both strategies could be used to synergistically enhance reprogramming efficiency.

In this study, we provide preliminary evidence of the pathways relevant for RIS. The up-regulation of p16\ ink4a \ is controlled by the epigenetic remodeling of the INK4a/ARF locus mediated, at least in part, by JMJD3. The p53/p21\ cip1 \ pathway is engaged at different levels in response to the expression of reprogramming factors. Similar to what is observed during replicative-induced senescence and OS, DNA damage is associated with the process. In addition, data extrapolated from the individual expression of reprogramming factors shows that the activation of p21\ cip1 \ seems a key endpoint in which different signals converge.

Reprogramming by introduction of defined factors started with empiric assays based in the knowledge of factors required for pluripotency [Takahashi and Yamanaka 2006]. Additional information about the molecular mechanism behind reprogramming can be used to add logic to the method. The study of developmental reprogramming of zygotes to pluripotent embryos can render us valuable clues. miRNAs up-regulated during that process are also required to sustain the rapid cell proliferation and cell cycle profiles associated with ES cells [Wang et al. 2008]. We showed that the expression of some of those miRNAs is also up-regulated in iPS but remains low at early times upon expression of reprogramming factors in fibroblasts. Ectopic expression of the miR-302 cluster alleviates RIS. Consistent with our hypothesis that inhibition of senescence enhances reprogramming efficiency, members of the miR-290 and miR-302 miRNA clusters have been shown to increase the efficiency of reprogramming [Judson et al. 2009]. Although the generation of iPS is an artificial process, this exemplifies how we can learn from its mechanisms and relate them to other physiologically relevant processes.

In summary, factors and processes involved in reprogramming evoke cellular senescence. Although the link between senescence and cancer makes impractical the derivation of iPS cells lacking tumor suppressors such as p53 or p16\ ink4a \, treating cells during reprogramming with reversible compounds to alleviate or inhibit senescence transiently could be a feasible alternative. Defining the precise timing in which temporary inhibition of senescence is needed to improve reprogramming will be the next logical step. Besides alternative approaches, maybe transient transfection with siRNAs targeting senescence effectors could be sufficient to improve the efficiency of reprogramming. Multiple groups are concentrating their efforts on finding strategies to enhance reprogramming efficiency. In addition to unbiased screening approaches, the knowledge of the reprogramming process has already suggested compounds or genetic elements that relax epigenetic regulation or that improve the cloning efficiency of ES cells as potential candidates to enhance reprogramming. We believe that senescence will be a key process to target in the pursuit of finding more efficient strategies for deriving iPS cells.

Materials and methods

Cell culture and retroviral infection

IMR90 and BJ human primary fibroblasts were from the American Type Culture Collection. MEFs knocked out for different genes were a gift from S. Lowe (Cold Spring Harbor Laboratory). Human and mouse fibroblasts were maintained in DMEM with 10% fetal calf serum (FCS). Methods used for isolation of MEFs and retrovirus production and infection have been described [Barradas et al. 2009].

Plasmids

Vectors encoding E6, E7, E6 + E7, H-RASV12G, or c-Myc have been described [Barradas et al. 2009]. The plasmid encoding miR-302 was generated by R. Agami [NKI, Amsterdam, The Netherlands] and obtained from Geneservice. Retroviral plasmids derived from pBABE puro encoding reprogramming factors are listed in Supplemental Table 1.

RNAi

Vectors encoding shRNAs targeting human p16\ ink4a \, human p53, and mouse Ink4a/Arf have been described previously [Barradas et al. 2009]. Sequences for shRNAs targeting human p21\ cip1 \ were cloned as double-stranded 21-mers in pRetroSuper, and are shown in Supplemental Table 2.
Antibodies
The antibodies used in this study are listed in Supplemental Table 3.

Immunofluorescence
Immunofluorescence was performed using an automated high-throughput microscope [InCell Analyzer 1000, GE]. Image processing and quantification was performed using InCell Investigator software (GE). A detailed description is given in the Supplemental Material.

BrdU, SA β-Gal assays, and crystal violet staining
BrdU labeling was performed for 16 h. SA β-Gal assays and crystal violet staining were performed as described previously (Barradas et al. 2009).

Quantitative RT–PCR analysis
Quantitative RT–PCR analysis was performed using standard protocols as described in Barradas et al. (2009). The primer sets and TaqMan 6-carboxyfluorescein (FAM) Gene Expression Assays [Applied Biosystems] used are listed in Supplemental Table 2 or in Pereira et al. (2008).

ChIP
Protocols, primers, and antibodies for ChIP have been described previously (Barradas et al. 2009).

Reprogramming of human and mouse fibroblasts
Reprogramming of human and mouse fibroblasts was carried out following standard protocols. A detailed description is given in the Supplemental Material.

Differentiation of hiPS cells
Differentiation of hiPS cells was carried out following protocols developed to differentiate hES cells. A detailed description is provided in the Supplemental Materials.

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