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Tumorigenicity is a well-documented risk to overcome for pluripotent or multipotent cell applications in regenerative medicine. To address the emerging demand for safe cell sources in tissue regeneration, we established a novel, protein-based reprogramming method that does not require genome integration or oncogene activation to yield multipotent fibromodulin (FMOD)-reprogrammed (FReP) cells from dermal fibroblasts. When compared with induced pluripotent stem cells (iPSCs), FReP cells exhibited a superior capability for bone and skeletal muscle regeneration with markedly less tumorigenic risk. Moreover, we showed that the decreased tumorigenicity of FReP cells was directly related to an upregulation of cyclin-dependent kinase inhibitor 2B (CDKN2B) expression during the FMOD reprogramming process. Indeed, sustained suppression of CDKN2B resulted in tumorigenic, pluripotent FReP cells that formed teratomas in vivo that were indistinguishable from iPSC-derived teratomas. These results highlight the pivotal role of CDKN2B in cell fate determination and tumorigenic regulation and reveal an alternative pluripotent/multipotent cell reprogramming strategy that solely uses FMOD protein.

Introduction

Local endogenous stem/progenitor cells are often unable to adequately reestablish tissue integrity and function after large-volume tissue loss from trauma, degenerative pathologies, or aging. In these cases, exogenous replacement cells derived from the stem or progenitor cells are necessary to restore lost tissue (1). Unfortunately, despite decades of investigation, isolation and/or generation of safe and readily available regenerative cell sources remain major challenges. In particular, the tumorigenicity risks are not only posed by teratomas formed by embryonic stem cells (ESCs) and induced pluripotent stem cells (iPSCs), but also include the tumor-supporting and/or formation potential of mesenchymal stem cells (MSCs) (2, 3), the main cell source for regenerative medicine used in the last few decades (4).

To address the emerging demand for safe and easily obtainable cell sources for tissue regeneration, we previously established an alternative strategy using continuous recombinant human fibromodulin (FMOD) stimulation under serum-free conditions to reprogram human dermal fibroblasts into multipotent cells (5). FMOD reprogramming shares several technological conveniences seen with iPSC generation. Both methodologies (a) offer freedom from the ethical and logistical constraints overshadowing ESC generation; (b) reprogram autologous cells, which significantly reduces the risk of immunogenic rejection; and (c) transform human dermal fibroblasts, a cell type that can be easily obtained and expanded from a skin biopsy (6), which may be preferred over the invasive harvest procedures required to obtain sufficient MSCs. Using this technology, dermal fibroblasts isolated from donors of different ages and sexes have been successfully reprogrammed into a multipotent state (5, 7). The resultant dome-shaped, clustered FMOD-reprogrammed (FReP) cells are easily separated from the surrounding spindle-shaped, monolayer FReP-basal cells with a xeno-free and enzyme-free reagent developed for passaging human pluripotent stem cells (7). Importantly, FReP cells exhibit pluripotency marker expression and triploblastic differentiation capabilities similar to those of ESCs and iPSCs (5). From an efficacy standpoint, we previously demonstrated that human FReP cells formed more bone than human iPSCs in a critical-sized SCID mouse calvarial defect model (7). Moreover, unlike iPSCs, implantation of FReP cells in the kidney capsule of immunodeficient mice did not result in teratoma or any other tumor formation (5).

After successfully using FReP cells to form bone, this current study focused on a more difficult challenge of regenerating skeletal muscle. Skeletal muscle constitutes 40%–50% of human body mass, and its significant loss can result in life-altering disabilities (8, 9). However, there are no readily available stem or progenitor cell sources for skeletal muscle repair (10–12). Moreover, the highly vascularized microenvironment of the muscle compartment can
be especially conducive to tumor formation — indeed, intramuscular implantation is a common route for teratoma formation when testing for cellular pluripotency (13). Thus, using nontumorigenic cells as starting materials is a key safety consideration for skeletal muscle regeneration.

Here, we demonstrate the persistence, engraftment, and myogenicity of FReP cells, without tumorigensis, in the tibialis anterior (TA) muscles of SCID mice. Moreover, FReP cell implantation into environments known to favor pluripotent cell teratoma formation, such as adult male Fox Chase SCID Beige mouse testes (13), did not result in tumor formation during the 4-month experimental period, which confirms the low tumorigenic potency of FReP cells. Furthermore, the FMOD-mediated reprogramming process itself is associated with marked upregulation of cyclin-dependent kinase inhibitor 2B (CDKN2B; also known as p15) (5) — an additional feature that helps minimize tumorigenicity — in FReP cells and potentially in other pluripotent/multipotent cells.

Results

FReP cells exhibit significant myogenic differentiation potential in vitro. In vitro, we used an established 2-stage skeletal myogenic differentiation protocol (5) that was developed for conventional retrovirus-mediated BJ fibroblast–derived iPSCs (BJ-iPSCs) (14) on FReP cells. In addition, human satellite cells were used as a positive control in vitro. The differentiated FReP cells expressed myogenic makers in a fashion similar to that of myogenically induced BJ-iPSCs, including myogenic differentiation 1 (MYOD1), sarcomeric α-actinin (ACTN), α-sarcomeric actin (ACTA1), myosin, and desmin (DES; Figure 1A). In contrast, unprogrammed parental BJ fibroblasts and FReP-basal cells failed to undergo myogenic differentiation when subjected to the same myogenic protocol (Figure 1A). Since gene profiling is a very reliable tool for analyzing the expression of a focused panel of genes, the “molecular blueprint” of FReP cell myogenic differentiation was comprehensively studied by a commercially available PCR array that revealed significant upregulation of 84 myogenesis-related genes in the myogenically stimulated FReP cells (Figure 1B and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI125015DS1). In addition, myogenically differentiated FReP cells expressed creatine kinase activity similar to that of human satellite cells (Figure 1C), indicating comparable in vitro functional myogenic potential between FReP cells and satellite cells (15, 16). Taken together, these data suggest that FReP cells are fully capable of behaving like muscle progenitor cells.

FReP cell implantation leads to skeletal muscle generation in vivo. To validate the myogenic potential of FReP cells in vivo, 5 × 10⁵ cells that had not undergone any form of premyogenic stimulation were implanted in the left TA muscles of 2-month-old male SCID mice. All negative controls — PBS vehicle (no cells), BJ fibroblasts, and FReP-basal cells — did not alter the TA muscle mass at 6 weeks after implantation (Figure 2A). Only limited numbers of BJ fibroblasts and FReP-basal cells survived in vivo. Surviving BJ fibroblasts were found on the surface of myofibers, while surviving FReP-basal cells were detected in some myofibers (Figure 2B). Meanwhile, retrovirus-mediated BJ-iPSCs, acting as a positive control, showed differentiation and engraftment that directly and significantly boosted muscle mass as evidenced by the spatial colocalization of human markers with the skeletal muscle markers (Figure 2B and Supplemental Figure 1). Excitingly, FReP cell implantation increased muscle mass to an even greater extent than retrovirus-mediated BJ-iPSC implantation (Figure 2A). Meanwhile, a broad spatial overlap of human markers with skeletal muscle markers confirmed the myogenic commitment and engraftment of FReP cells in the SCID mouse TA muscles (Figure 2B and Supplemental Figure 1). Overall, FReP cells exhibited superior skeletal muscle generation in vivo when compared with iPSCs.

FReP cells have less tumorigenic potential than iPSCs. Notably, 2 of 8 animals (25%) that underwent implantation of retrovirus-mediated BJ-iPSCs into their uninjured TA muscles experienced tumor formation with active cell proliferation instead of skeletal muscle generation (Figure 2A and Supplemental Figure 2). Neither FReP-basal nor FReP cell implantation led to tumor formation during skeletal muscle (Figure 2) or bone (5, 7) regeneration, suggesting less tumorigenic potential than iPSCs.

Since iPSC tumorigenesis is considered to be driven by mutations associated with uncontrollable proliferation (17, 18), cellular proliferation was examined. In agreement with previous studies (5, 6), retrovirus-mediated BJ-iPSCs exhibited extremely rapid proliferation, while FReP cells proliferated minimally under undifferentiated conditions in vitro (Figure 3A). Next, a soft agar colony formation assay, the standard tumorigenicity test, was used to examine anchorage-independent cellular survival under a low-nutrient and -oxygen microenvironment (19). After 14 days of cultivation with 10 μM Y-27632, the survival of BJ fibroblasts was negligible, while retrovirus-mediated BJ-iPSCs actively proliferated and formed colonies (Figure 3, B and C). Neither FReP-basal nor FReP cells proliferated or formed colonies; however, FReP-basal cells largely adopted a spindle shape while FReP cells remained morphologically round in the soft agar (Figure 3, B and C).

It is worth noting that the correlation between the soft agar colony formation assay and in vivo tumorigenicity tests is imprecise (20). The soft agar colony formation assay may underestimate the tumorigenic potential of reprogrammed pluripotent/multipotent cells, because only a small portion of tumorigenic iPSCs form colonies in the gel, owing to the dissociation-induced apoptosis of potentially tumorigenic iPSCs when conducting the assay (19, 21, 22). Since the intratesticular stromal cells may produce a milieu that is more supportive for implanted cells than subcutaneous or intramuscular microenvironments (13), 1 × 10⁴ cells with 30 μL of Matrigel carrier were intratesticularly implanted into Fox Chase SCID Beige mice to further assess the tumorigenicity of FReP cells in vivo. In this system (20), intratesticular implantation of retrovirus-mediated BJ-iPSCs resulted in 100% (10/10) of the implanted animals developing teratomas with progressive growth (Figure 4A and Supplemental Figure 3B). Excitingly, none of the animals implanted with FReP or FReP-basal cells formed teratomas during the entire 4-month experimental period. Meanwhile, no human markers were detected in mouse testes implanted with FReP cells at 4 months after implantation (Figure 4A). A similar lack of human marker expression was also noted after BJ fibroblast or FReP-basal cell implantation in mice at
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Figure 1. FReP cells display myogenic differentiation potential in vitro. (A) Myogenic markers ACTN, MYOD, ACTA1, myogenin, and desmin were found in FReP cells after cultivation using the established in vitro 2-stage skeletal myogenic differentiation protocol. Under the same differentiation conditions, parental unprogrammed BJ fibroblasts and FReP basal cells were used as negative controls, while retrovirus-mediated BJ iPSCs and differentiated human satellite cells were used as positive controls. White arrowheads indicate the fusing myogenically differentiated cells; scale bar: 100 μm. (B) A PCR array revealed significantly upregulated myogenesis-related genes in FReP cells during the 3-week in vitro myogenic differentiation period. Unprocessed original Ct data are shown in Supplemental Table 1. n = 3 independent experiments. (C) The creatine kinase activity assay suggests that, after undergoing myogenic differentiation, FReP cells have biological activities similar to those of human satellite cells in vitro. *P < 0.005 (analyzed by 1-tailed Mann-Whitney and Kruskal-Wallis ANOVA tests); n = 6.

The same time (Figure 4A). This suggests that, like BJ fibroblasts, FReP basal and FReP cells that did not undergo predifferentiation stimulation did not spontaneously engraft or differentiate into the host testis tissue. Our previous kidney capsule injection (5) and current intramuscular and intratesticular implantation studies collectively constitute strong evidence that FReP cells are less likely to generate tumors than iPSCs.

FReP cells and iPSCs exhibit different proto-oncogene and tumor suppressor gene expressions. Since FMD reprogramming is a genome integration-free technology (5, 7), the global gene profile of BJ-derived FReP cells (GEO accession number GSE104830) was first compared with that of BJ-derived iPSCs generated through non–genome integration procedures (GEO accession numbers SRR500985/6/7/8, ref. 23; and SRR1583694/5, ref. 24) to gain insight into the mechanisms governing the disparate tumorigenic natures of FReP cells and iPSCs. A total of 2300 highly differentially expressed mRNAs (fold change ≥ 2), identified by the TopHat-Cufflinks package (25) through the Galaxy platform (26), were further analyzed by the DAVID Bioinformatics Resource 6.8 for functional annotation (27). This analysis revealed an enrichment of genes involved in the “pathways in cancer” (Supplemental Figure 4A) with significant similarity (κ = 1.0). The differential expression of genes that aligned with human proto-oncogenes and tumor suppressor genes (TSGs) recognized in the UniProt database (28) is displayed graphically (Supplemental Figure 4, B and C).

In comparison with non–genome integration BJ-iPSCs, FReP cells displayed more proto-oncogenes with relatively lower expression levels (Supplemental Figure 4B). Meanwhile, in comparison with retrovirus-mediated BJ-iPSCs used in this study, quantitative reverse transcriptase PCR (qRT-PCR) confirmed that both FReP cells and FReP basal cells have significantly lower expression levels of proto-oncogenes (Figure 4B) such as MYC, known to markedly accelerate cell proliferation and promote iPSC generation and tumor formation (29), and lin-28 homolog A (LIN28), documented to augment iPSC induction efficiency in a cell division-rate–dependent manner (30). Meanwhile, when compared with non–genome integration BJ-iPSCs, FReP cells have a greater number of TSGs with higher expression levels (Supplemental Figure 4C).

Further analyses of highly expressed TSGs in FReP cells using the STRING database to retrieve protein–protein interactions (31) emphasized the central roles of tumor protein p53 (TP53 in humans) and CDKN2B (Supplemental Figure 4D).

Accumulating evidence demonstrates that activating TSGs, particularly TP53, cyclin-dependent kinase inhibitor 1A (CDKN1A, also known as p21, which produces a translational product that mediates TP53-induced cell cycle arrest and tumor suppression; ref. 32), CDKN2A (also known as p16), and CDKN2B, reduces the efficiency of iPSC generation and limits the cancerous transformation of iPSCs predominantly by inhibiting cell division (17–19, 30, 33–37). For example, Menendez et al. reported that iPSCs derived from mouse embryonic fibroblasts (MEFs) with an elevated level of p53 (38) or the Ink4a/Arf locus (which encodes Cdkn2b, Cdkn2a, and Arf [an alternative transcript of Cdkn2a] (39) have less tumorigenic potential than iPSCs derived from wild-type MEFs (19). In comparison with parental BJ fibroblasts, retrovirally derived BJ-iPSCs also have significantly less TP53 expression (Figure 5A). As seen in the MEF-derived iPSC study (40), expression of CDKN1A and CDKN2B, but not CDKN2A, was markedly reduced in retrovirus-mediated BJ-iPSCs (Figure 5, B–D). Meanwhile, these 4 genes were all upregulated in both FReP basal and FReP cells (Figure 5), which aligns with our previous report in which these 2 populations were not distinguished from one another (5). Notably, higher expression levels of CDKN1A and CDKN2A were observed in FReP basal cells than in FReP cells (Figure 5, B and C), while a more robust increase in CDKN2B was detected in FReP cells (Figure 5D).

FReP cells reprogrammed from CDKN2B-knockdown fibroblasts acquire teratoma-forming, pluripotent characteristics. To further investigate the effects of TP53 and these CDKNs on the proliferation, multipotency, and tumorigenesis of FReP cells, stable TP53- or CDKN2B-targeted (KD) BJ cells were established, respectively, in which the RNAi effects were extended beyond the FMD reprogramming process (Figure 6A and Supplemental Figure 5). All of the KD BJ cells had reduced CDKNIA levels, while CDKN2A KD did not affect the expression of TP53, CDKN2A, or CDKN2B. Also, because of the structure of the INFA4/ARF locus, expression of CDKN2A and CDKN2B was decreased by some degree in response to both CDKN2A and CDKN2B KD (Supplemental Figure 5). The upregulation of CDKN2B induced by FMD reprogramming was also blocked by CDKN2B KD: CDKN2B expression increased 21.4-fold in the scrambled shRNA-transfected BJ fibroblasts (scramble FReP cells) in response to FMD reprogramming; but in CDKN2B-KD BJ fibroblasts, CDKN2B transcription was not altered by continuous FMD treatment (Figure 6A). Thus, the expression of CDKN2B in FReP cells derived from CDKN2B-KD BJ fibroblasts (CDKN2B-KD FReP cells) was comparable to that seen in BJ-iPSCs.

Although CDKN2B-KD FReP cells presented higher expression levels of POUSF1 (also known as OCT4) in comparison with scramble FReP cells or the aforementioned FReP cells derived from BJ fibroblasts without shRNA transfection (in brief, FReP cells; Supplemental Figure 6A), all of these 3 FReP cells had similar multipotent differentiation capabilities and, importantly, myogenic differentiation potentials in vitro (Figure 6, B and C, and Supplemental Figure 6B). To rule out the possibility that the retained multipotent differentiation capability of CDKN2B-KD FReP cells was due to inefficient CDKN2B downregulation at the protein level, we carried out Western blotting to further confirm the markedly decreased expression of CDKN2B protein in...
CDKN2B-KD FReP cells (Supplemental Figure 7). This showed that decreased CDKN2B expression does not diminish the multipotency of FReP cells. Conversely, in agreement with a previous report affirming that permanent p53-p21 pathway suppression impairs the maintenance of iPSC identity (36), TP53- and CDKN1A-KD FReP cells exhibited reduced expressions of all the pluripotent markers, except for NANOG (Supplemental Figure 6A), and impaired myogenic differentiation (Figure 6B) when compared with scramble FReP cells and CDKN2B-KD FReP cells. Surprisingly, TP53, CDKN1A, and CDKN2A KD reduced the multilineage differentiation capacity of FReP cells (Supplemental Figure 6B), suggesting that the upregulation of TP53 and CDKNs is required for FMOD reprogramming and endows FReP cells with their unique triploblastic differentiation potency.

Meanwhile, all tested TP53- or CDKN-KD FReP cells exhibited increased proliferation rates in comparison with FReP cells or scramble FReP cells (Figure 7A). However, in vitro assays demonstrated that only CDKN2B-KD FReP cells were able to form colonies in the soft agar, similar to observations in retrovirus-mediated BJ-iPSCs (Figure 7B and C). Moreover, in comparison with FReP cells and scramble FReP cells, CDKN2B-KD FReP cells had significantly elevated expressions of the aforementioned proto-oncogenes, except for DEK (Figure 7D). Intratesticular implantation of CDKN2B-KD FReP cells, but not the other aforementioned knockdown FReP cells, resulted in teratoma formation in 100% (10/10) of the tested animals (Figure 8A and Supplemental Figure 8E). The CDKN2B-KD FReP cell-derived teratomas were indistinguishable from retrovirus-mediated BJ-iPSC-derived teratomas (Figure 8B). These data constitute the evidence that CDKN2B KD alone converted nontumorigenic, multipotent FReP cells to teratoma-forming pluripotent stem cells.

Discussion

To address the shortage of tissue-specific progenitor cells, multipotent and pluripotent cells offer exciting alternatives for cell-based tissue repair (4). However, safety concerns,
particularly tumorigenicity, must be thoroughly surmounted before the widespread usage of multipotent and pluripotent cells in regenerative medicine (41).

Within the last few decades, MSCs have been widely investigated for their potential use in tissue regeneration because of their multipotent and immunomodulatory properties (42), even though they are generally harvested through more painful and invasive procedures than simple skin biopsy, such as muscle tissue biopsy (43), bone marrow aspiration (44), and liposuction (45). In addition, it is now well documented that MSCs exhibit transient and surprisingly low (less than 1%) engraftment in newly formed tissues (46, 47). Thus, MSCs assert their regenerative potency by secreting trophic factors instead of directly differentiating into the target tissues (48–52). As an off-target effect, various cytokines, chemokines, and growth factors secreted by MSCs are known to increase the proliferation, migration, and angiogenesis of tumor cells (2).

Human MSCs can develop chromosomal aberrations during cultivation (53), undergo spontaneous tumorigenic transformation (54), promote the growth of cocultured glioblastoma cells in vitro, and support glioblastoma development in vivo (3). Animal studies also demonstrate the direct and/or indirect involvement of MSCs in sarcomagenesis (2), especially with inflammatory stimulation (55) that typically arises in injury scenarios. Thus, tumorigenesis is recognized as another risk of using MSCs in humans.

The breakthrough discovery of iPSCs created enthusiasm to use these pluripotent cells for tissue regeneration. However, the widely accepted procedure for iPSC generation, in which transcription factors essential for embryonic development (such as the Yamanaka or Thomson factors) are introduced into the genome of target somatic cells, is known to induce unwanted gene activation and genomic alterations that pose considerable hazards for clinical use (56). For example, in the first clinical trial using iPSC-derived retinal pigment epithelial (RPE) cells to replace age-related macular degeneration RPE cells, 3 DNA deletions were detected in the autologous iPSCs and their derived RPE cells in 1 of the 2 patients (50%), which halted the planned transplantations (57). Moreover, retrovirus-mediated and transgene-free human iPSCs exhibit similar tumorigenicity with no appreciable difference in teratoma formation capability or teratoma microvascular density (17, 37). Although profound efforts are devoted to replacing transcription factors tightly associated with tumor progression with defined chemicals, current research demonstrated that BrdU, a mutation inducer that can incorporate into newly synthesized DNA by replacing thymidine during DNA replication, is absolutely required for achieving chemical induction of pluripotency (58). Considering that iPSCs also possess a potential risk for somatic tumor development that is not present with ESCs (17), the tumorigenic risk of iPSCs remains a credible concern regarding their use clinically.

It has been hypothesized that the pluripotent cell microenvironment plays important roles determining cell fate, as well as the maintenance and induction of pluripotency/multipotency (59). In support of this, early pioneering studies demonstrated the success-

**Figure 3. FReP cells proliferate minimally in undifferentiating conditions and do not form colonies in a soft agar colony formation assay.** (A) A cell proliferation assay was carried out in 96-well culture plates after 2000 cells per well were cultured in undifferentiating conditions for 3 days. (B and C) A soft agar colony formation assay was performed after 14 days of cultivation. Five thousand cells per well were initially seeded. Data are presented as mean values. Gray dashed lines indicate the original cell seeding densities; black asterisks indicate significance in comparison with BJ fibroblasts; blue asterisks indicate significance in comparison with FReP cells. **P < 0.005** (A and B; analyzed by 1-way ANOVA and 1-tailed 2-sample t tests); n = 6; scale bar: 500 μm (C).
Figure 4. Intratesticular implantation of FReP cells does not lead to tumorigenesis in Fox Chase SCID Beige mouse testes. (A) Gross appearance and histological evaluation (H&E staining) of adult Fox Chase SCID Beige mouse testes that were intratesticularly implanted with $1 \times 10^6$ cells were documented at 4 months after implantation. All implanted mice testes are shown in Supplemental Figure 3 (10 mice per group). In addition, by tracking of human mitochondria in vivo, significant survival of the implanted human cells was only observed in the BJ-iPSC group, in which teratoma formation was also detected. Scale bars: 5 mm (black), 1 mm (blue), or 100 μm (yellow). (B) The expressions of multiple proto-oncogenes were compared between parental BJ fibroblasts, retrovirus-mediated BJ-iPSCs, FReP-basal cells, and FReP cells. ERBB3, erb-b2 receptor tyrosine kinase 3; DEK, DEK proto-oncogene; DNMT3B, DNA methyltransferase 3β; FLT3, fms-related tyrosine kinase 3; FOXO1, forkhead box 1; LIN28, lin-28 homolog A; KIT, KIT proto-oncogene receptor tyrosine kinase; POU2F1, POU class 2 homeobox 1; TDGF1, teratocarcinoma-derived growth factor 1. Data are normalized to those of the BJ fibroblasts and presented as mean ± SD. **P < 0.005 (analyzed by 1-way ANOVA and 1-tailed 2-sample t tests); n = 3 independent experiments performed in duplicate. Dashed lines indicate the gene expression levels of BJ fibroblasts; black asterisks indicate significance in comparison with BJ fibroblasts; blue asterisks indicate significance in comparison with FReP cells.
ful induction of multipotent stem cells from somatic cells using extracts from *Xenopus* eggs (60), fish oocytes (61), ESCs (62), and even carcinoma cells (62). Interestingly, recent studies show that fetal MSCs, such as umbilical cord–derived MSCs (UCMSCs), express both specific ESC and adult MSC markers, suggesting that UCMSCs may represent an intermediate cell stage that is developmentally between the adult MSCs and ESCs (63–65). It is of interest that UCMSCs are predominantly harvested from Wharton’s jelly, a proteoglycan-rich connective tissue (66, 67); this suggests that recreating a proteoglycan-rich microenvironment may facilitate or induce somatic cell reprogramming to some degree. FMOD exposure may reconstitute a specific microenvironment resembling quiescent stem cells in multiple ways (5), continuous stimulation by FMOD in serum-free conditions is essential for establishing and/or maintaining the fetal and stem cell niches (71, 72). These collective data suggest that FMOD is a critical component for the maintenance of endogenous stem cell niches. To test this hypothesis, we cultured human newborn foreskin BJ fibroblasts and normal adult dermal fibroblasts in FMOD and were able to induce successful reprogramming of these cell populations (5). Excitingly, our previous (7) and current studies demonstrate that the FMOD-reprogrammed FReP cells exhibit a superior capability for bone and skeletal muscle regeneration in comparison with iPSCs. More importantly, the implantation of FReP cells, even in an undifferentiated state, does not cause teratoma or any other kind of tumor formation in vivo in all tests to date. Therefore, FReP cells are a readily available and potentially safer source for musculoskeletal system regeneration as shown here and potentially other regenerative medicine applications.

The decreased tumorigenic potential and proliferation activity, as well as the favorable oncogene and TSG expression profile, also highlight that both the FMOD reprogramming process and the resultant FReP cells are fundamentally distinct from iPSCs. Interestingly, FReP cells bear several critical characteristics of multilineage-differentiating stress-enduring (MUSE) cells (5, 73): they (a) express pluripotent markers, albeit at relatively lower levels than ESCs and iPSCs; (b) can differentiate into all 3 germ-line cell lines under specific induction conditions; (c) have low levels of proto-oncogenes, such as *LIN28*; (d) retain a stable karyotype; and, most importantly, (e) do not form teratomas. Although FReP cells and MUSE cells are both excluded from being considered pluripotent by the stringent mandatory criteria of teratoma formation in vivo, they may represent a distinct group of cells with a triploblastic differentiation capability that holds tremendous potential in regenerative medicine.

However, FReP cell generation is distinct from MUSE cell collection. Activation and isolation of MUSE cells require severe cellular stress conditions, such as lengthy incubation and digestion, hypoxia, and low temperatures (73), which assist in killing all other viable cells. Conversely, FMOD reprogramming does not require hypoxia or low temperatures, and the resultant FReP cells and FReP-basal cells are both viable. Given the fact that FReP cells resemble quiescent stem cells in multiple ways (5), continuous FMOD exposure may reconstitute a specific microenvironment similar to the niche of quiescent stem cells, and thus induce reprogramming of somatic cells.

Reprogramming and tumorigenesis appear interconnected in many reports examining induced pluripotency (18), although the mechanism governing the transition from nontumorigenic somatic cells to tumorigenic iPSCs remains an enigma. Nevertheless, our current studies demonstrate that FReP cells derived from *CDKN2B*-KD fibroblasts acquired undesirable tumorigenic characteristics. Still, to our knowledge, this represents the first report of conversion of nontumorigenic, multipotent cells obtained via a non-genome integration procedure into a teratoma-forming, pluripotent state simply by suppression of the upregulation of *CDKN2B*. These results indicate that, in addition to the conventional transcriptional factor–based procedure, continuous stimulation by FMOD in serum-free conditions while blocking the increase in *CDKN2B* may be an alternative
strategy to convert somatic cells to pluripotent states. These results highlight the essential role of CDKN2B in governing the transition of multipotent to pluripotent cells, especially during reprogramming procedures without genome integration or oncogene activation.

CDKN2B abnormalities have been associated with many forms of carcinogenesis, including leukemias, carcinomas, and melanomas. As an effector of TGFβ, CDKN2B inhibits cyclin-dependent kinase 4/6, and thus induces a G1-phase cell cycle arrest (74). However, since most previous studies focused on

Figure 6. TP53 and CDKN gene–KD FReP cells have different gene expression profiles and myogenic differentiation potentials. (A) Gene expression of TP53 and CDKN genes was assessed in the FReP cells derived from different KD BJ fibroblasts. Data are presented as mean ± SD and normalized to those of the BJ fibroblasts without any shRNA transfection. *P < 0.05, **P < 0.005 (analyzed by 1-way ANOVA and 1-tailed 2-sample t tests); n = 3 independent experiments performed in duplicate. Black dashed lines indicate the gene expression levels of BJ fibroblasts without any shRNA transfection (in brief, BJ fibroblasts); red dashed lines indicate the gene expression levels of BJ-iPSCs; blue dashed lines indicate the gene expression levels of FReP cells derived from BJ fibroblasts without any shRNA transfection (in brief, FReP cells); gray asterisks indicate significance in comparison with FReP cells generated from scrambled shRNA–transfected BJ fibroblasts (scramble FReP cells). (B and C) Myogenic differentiation of KD FReP cells was assessed by myogenic marker staining (B) and creatine kinase activity assay (C) in vitro. White arrowheads indicate the fusing myogenic differentiated cells. Scale bar: 100 μm (B). **P < 0.005 (analyzed by 1-tailed Mann-Whitney and Kruskal-Wallis ANOVA tests, C); n = 6.
Figure 7. TP53 and CDKN gene KD diversely alters FReP cell proliferation, anchorage-independent survival ability in soft agar, and proto-oncogene expression in vitro. 

(A) A cell proliferation assay was carried out in 96-well culture plates after 2000 cells per well were cultured in undifferentiating conditions for 3 days. 

(B and C) A soft agar colony formation assay was performed after 14 days of cultivation. Five thousand cells per well were seeded initially. 

(D) The expressions of multiple proto-oncogenes in scramble FReP cells and CDKN2B-KD FReP cells were compared. Data are presented as mean values (A and B) or mean values normalized with those of the BJ fibroblasts ± SD (D). **P < 0.005 (analyzed by 1-way ANOVA and 1-tailed 2-sample t tests); n = 6 (A–C) or 3 independent experiments performed in duplicate (D). Gray dashed lines indicate the original cell seeding densities (A and B); black dashed lines indicate the gene expression levels of BJ fibroblasts (D); red dashed lines indicate the gene expression levels of BJ-iPSCs (D); blue dashed lines indicate the cell densities (A and B) or the gene expression levels (D) of FReP cells, respectively; gray asterisks indicate significance in comparison with scramble FReP cells. Scale bar: 500 μm.
the INK4/ARF locus as a whole without separating CDKN2B from CDKN2A, these studies did not fully elucidate the isolated function of CDKN2B. Hence, the contribution of low CDKN2B levels to tumorigenesis may be underestimated. Taking a closer look at the INK4/ARF locus, CDKN2A seems to play a more vigorous role in oncogenesis than CDKN2B. For instance, in acute lymphoblastic leukemia, CDKN2A is often deleted without CDKN2B deletion, while a sole CDKN2B deletion is almost never detected (75). However, our current results demonstrate that sustained suppression of CDKN2B, instead of CDKN2A, during FMOD reprogramming plays the predominant role in introducing a tumorigenic potential into the nontumorigenic FReP cells. This suggests that CDKN2A and CDKN2B may have different bioactivities and that further targeted studies on the role of CDKN2B in reprogramming, maintenance, and tumorigenesis of multipotent/pluripotent cells, including MUSE cells and iPSCs, are warranted.

Moreover, on the basis of the clear, inverse relationship between tumorigenesis and CDKN2B levels observed in this study, elevating CDKN2B expression may be an interesting, alternative antitumor strategy to decrease the tumorigenic risk of the therapies based on pluripotent/multipotent cells (such as iPSCs). However, it is worth noting that iPSCs exhibit significantly lower INK4/ARF locus expression than their parental fibroblasts (76). In particular, CDKN2B is more severely repressed during iPSC reprogramming than CDKN2A and ARF (76). Meanwhile, transient silencing of the INK4/ARF locus is sufficient to accelerate the reprogramming progress and increase the number of iPSCs (76). Since INK4/ARF locus inhibition promotes iPSC generation, it is highly possible that forcing CDKN2B expression may fundamentally impede the reprogramming process that yields iPSCs. An alternative strategy to test whether a CDKN2B-based antitumor strategy would work in iPSCs is to consistently upregulate CDKN2B expression in fully reprogrammed iPSCs to suppress their tumorigenic potency. However, despite the low efficiency, the potential off-target effects of human iPSC genome editing often lead to high cell mortality and, more importantly, introduce mutations that are independent of the gene of interest, and thus confound the analysis of the tumorigenic risk of the genome-edited cells (77-82). Even more crucially, because of the cluster-forming nature of iPSCs, it would be a major challenge to purify CDKN2B-overexpressing iPSCs from the surrounding iPSCs without CDKN2B upregulation, especially since there is no known surface marker that could assist this process currently. Forcing CDKN2B expression in reprogrammed iPSCs may also not be a practical and reliable way to reduce the tumorigenic risk of iPSCs, as even only a small portion of tumorigenic iPSC contamination can induce teratoma formation (83). Therefore, aside from understanding whether CDKN2B can prevent iPSC tumorigenesis, how to reliably evaluate and effectively implement the CDKN2B overexpression strategy in iPSCs remains a key challenge for future studies.

In summary, we demonstrated that FReP cells exhibit a superior capability for skeletal muscle regeneration with markedly less tumorigenic risk when compared with iPSCs. As FReP cells are fundamentally distinct from iPSCs, FReP cells may represent another class of cells with a triploblastic differentiation capability rather than teratoma-forming pluripotency that may have greater applications in regenerative medicine. Concomitantly, we identified that decreased FReP cell tumorigenicity is directly related to CDKN2B upregulation during the FMOD reprogramming process, highlighting an essential role for CDKN2B in regulating cell fate and tumorigenesis.

Methods

Cell culture. Human newborn foreskin BJ fibroblasts (ATCC CRL-2522; ATCC) were cultured in a 4:1 mixture of DMEM (containing 4 mM l-glutamine, 1.0 g/L glucose, and 1.5 g/L sodium bicarbonate; Thermo Fisher Scientific) and Medium 199 (Thermo Fisher Scientific), supplemented with 10% FBS (Thermo Fisher Scientific) and 1% penicillin/streptomycin (P/S; Thermo Fisher Scientific) (7). BJ fibroblasts were authenticated by Laragen Inc. and tested negative for mycoplasma contamination using the Universal Mycoplasma Detection Kit (ATCC; Supplemental Figure 9). BJ-iPSCs obtained by conventional retrovirus-mediated methods (14) were maintained on plates precoated with Matrigel human ESC-qualified (hESC-qualified) Matrix (BD Biosciences) with mTESR1 medium (STEMCELL Technologies) (7).

Human satellite cells were purchased from Lifeline Cell Technology and cultured according to the manufacturer’s instructions. After 1 week of culture in MyoLife Complete Myogenesis Differentiation Medium (Lifeline Cell Technology), the differentiated human satellite cells were used as a positive control for in vitro myogenic differentiation.

FMOD production. cDNA of a human FMOD transcript (GenBank NM_002023) was subcloned into a commercially available vector, pSecTag2A (Thermo Fisher Scientific), with a C-terminal His tag, and then transfected into CHO-K1 cells (ATCC) (84). After establishment of a stable FMOD expression clone, FMOD was produced and purified by a contract research organization, GenScript. Briefly, a stable human recombinant FMOD–expressing CHO-K1 cell line was cultured in 1 L of serum-free Freestyle CHO Expression Medium (Thermo Fisher Scientific) at 37°C with 5% CO2 in an Erlenmeyer flask. The cell culture supernatant was harvested after 10 days for purification with a HiTrap IMAC HP 1-mL column (GE Healthcare). The fractions from a 100-mM imidazole elution were collected and dialyzed against 20 mM PBS, pH 7.4. Next, the sample with low conductivity was loaded onto a HiTrap Q HP 1-mL column (GE Healthcare) for further purification. FMOD was then purified under nonreducing conditions, dia lyzed again (7), and then subjected to lyophilization. The purity of the FMOD product was 85%. FMOD was reconstituted in PBS and then underwent sterilization through a 0.22-μm filter (Thermo Fisher Scientific) before use (70).

FMOD reprogramming. BJ fibroblast cells (4 × 10^6 per well) were seeded in 6-well culture plates overnight to confluence and then exposed to 0.4 mg/mL recombinant human FMOD in DMEM supplemented with 1% P/S for reprogramming under serum-free conditions. The medium was exchanged with fresh medium daily (5, 7). After 21 days of continuous FMOD treatment, FReP cells were harvested by ReLeSR (an enzyme-free hESC and hiPSC selection andpassaging reagent; STEMCELL Technologies), which significantly increased the yield of FReP cells in comparison with a traditional manual scraping method (Supplemental Figure 10). The yielded cells were then cultured on Matrigel hESC-qualified Matrix–coated (BD Biosciences) plates with mTESR1 medium (STEMCELL Technologies) (7).

RNAi. Following the manufacturer’s instructions, plasmids harboring TP53 shRNA, CDKN1A shRNA, CDKN2A shRNA, or CDKN2B shRNA (Santa Cruz Biotechnology) were used to knock down the
expression of the respective target genes. Control shRNA Plasmid-A (Santa Cruz Biotechnology), which encoded a scrambled shRNA sequence, was used as a negative control for RNAi. For each target gene, the colony with the lowest gene expression of the target gene was selected from 5 candidate knockdown colonies for further investigation.

**Proliferation assay.** Two thousand cells per well were seeded into 96-well culture plates for a proliferation assay. After 3 days of incubation, cell proliferation was assessed by the Vybrant MTT Cell Proliferation Assay Kit (Molecular Probes) with an Epoch Microplate Spectrophotometer coupled with Gen5 software (version 2.0.4.11, BioTek Instruments Inc.).

**Soft agar colony formation assay.** Soft agar colony formation assay was performed with the CytoSelect 96-Well In Vitro Tumor Sensitivity Assay (Cell Biolabs Inc.) according to the manufacturer’s instructions. Five thousand cells per well were initially seeded with the supplied medium and 10 mM Rho-associated protein kinase (ROCK) inhibitor Y-27632 (STEMCELL Technologies). After 14 days of cultivation, representative photographs were captured on an Olympus IX71 microscope coupled with a DP73 camera and cellSens Standard 1.9 software (Olympus). Cellular anchorage-independent growth was quantified according to the instructions of the CytoSelect 96-Well In Vitro Tumor Sensitivity Assay with an Epoch Microplate Spectrophotometer.

**In vitro 2-stage skeletal myogenic differentiation.** Cells were seeded on Attachment Factor Protein–coated plates with myogenic medium I (DMEM supplemented with 10% FBS, 1% P/S, 10% horse serum [HS; Thermo Fisher Scientific], and 1% chicken embryo extract [CEE; Gemini Bio Products]) for 7 days. Then, myogenic medium II (DMEM supplemented with 1% FBS, 1% P/S, 1% HS, and 0.5% CEE) was used for another 14 days of cultivation. Half of the medium was renewed every 4 days (5, 85).

Antibodies against ACTA1 (catalog ab28052, clone Alpha Sr-1, Abcam), ACTN (catalog ab9465, clone EA-53, Abcam), desmin (catalog ab8592, Abcam), MYOD (catalog ab64159, Abcam), and myosin (catalog MAB1628, clone NOQ7.4D, MilliporeSigma) were used to confirm skeletal myogenic differentiation. All immunostaining, in terms of concentration, antigen retrieval, etc., was conducted based on the instructions of the manufacturers.

One microgram total protein of lysate from cells that underwent myogenic differentiation in vitro was used for a creatine kinase activity test (15, 16) using the Creatine Kinase Activity Assay Kit (MilliporeSigma) according to the manufacturer’s protocol. The test was performed at 37°C, and results were documented with an Epoch Microplate Spectrophotometer.

**In vitro neurogenic differentiation.** Cells were seeded on AggreWell 800 Plates with AggreWell medium (STEMCELL Technologies) for embryoid body (EB) formation, following the manufacturer’s instructions (5, 7). EBs were then transferred to a 6-well Ultra Low Cluster Plate (Costar, Corning Inc.) with KnockOut DMEM (Thermo Fisher Scientific) supplemented with 10% KnockOut Serum Replacement (Thermo Fisher Scientific), 2 mM L-glutamate (Thermo Fisher Scientific), 1% P/S, 10 μM all-trans retinoic acid (RA; Santa Cruz Biotechnology), and 100 nM of the N-terminal active fragment of human sonic hedgehog (Shh; R&D Systems) to generate spheres. Fresh RA was added every day, and the medium and supplements, including Shh, were replaced every 3 days. After 8 days of suspension culture, the induced spheres were transferred onto poly-ornithine/fibronectin-coated (MilliporeSigma) plates with DMEM/F12 medium (Thermo Fisher Scientific) supplemented with 2% FBS, 1% P/S, N2 Supplement (Thermo Fisher Scientific), 20 ng/mL glial-derived neurotrophic factor (Thermo Fisher Scientific), 20 ng/mL brain-derived neurotrophic factor (Thermo Fisher Scientific), 20 ng/mL ciliary neurotrophic factor (Thermo Fisher Scientific), and 1× B27 serum-free supplement (Thermo Fisher Scientific) (5, 86, 87). The medium was changed every 3 days. After 9 days, cells were fixed for immunocytochemistry examination with antibodies against neuron-specific βIII-tubulin (also known as TUJ1; catalog ab18207, Abcam) (5, 7).

**In vitro osteogenic differentiation.** Cells were seeded on Attachment Factor Protein–coated (Thermo Fisher Scientific) plates and cultured with MEM Alpha (Thermo Fisher Scientific) supplemented with 10% FBS, 50 μg/mL ascorbic acid (MilliporeSigma), 10 mM β-glycerophosphate (MilliporeSigma), and 1% P/S for 28 days. The medium was changed every 3 days. Mineralization was detected by alizarin red staining (LifeLine Cell Technology) (5, 7).

**In vitro pancreatic differentiation.** Cells were cultured in RPMI 1640 medium (Thermo Fisher Scientific) supplemented with 2% FBS, 2 mM L-glutamate, 1% P/S, and 100 ng/mL recombinant activin A (Thermo Fisher Scientific) for 4 days to differentiate into an endoderm derivative. Cells were cultured for another 16 days without activin A (5, 88, 89) before characterization with antibodies against Nkx6.1 (catalog F55A10, clone F55A10, Developmental Studies Hybridoma Bank), insulin (catalog 4590, Cell Signaling Technology), pancreatic and duodenal homeobox 1 (PDX1; catalog ab47267, Abcam), and amylin (catalog GTX11022, clone R10/99, GeneTex).

**Immunocytochemistry.** Samples selected for immunocytochemistry staining were fixed in prechilled acetone. DAPI (MilliporeSigma) was used for nuclear staining (5, 7). The StemLite Pluripotency Kit (catalog 9656, Cell Signaling Technology) was used to detect a panel of proteins that are specifically expressed in human pluripotent cells (5). Images were collected with the Leica TCS SP8 Confocal Laser Scanning Platform (Leica).

**RNA isolation.** RNA was extracted using the RNaseasy Mini Kit (Qiagen) with DNase (Qiagen) treatment to ensure that the samples were not contaminated with genomic DNA. RNA purity was assessed by the Epoch Microplate Spectrophotometer.

**PCR array quantitative RT-PCR.** PCR arrays are one of the most reliable tools for analyzing the expression of a focused panel of genes. To further establish the feasibility of FReP cells as an alternative safe cell source for skeletal muscle regeneration, we used PCR arrays to generate a “molecular blueprint” of FReP cell activities during myogenic differentiation. According to the manufacturer’s instruction, the RT2 Profiler PCR Array for Human Skeletal Muscle (Qiagen) was used to track the expression of myogenic-related genes. Three different cDNA templates were tested.

**qRT-PCR.** For qRT-PCR, 1.0 μg RNA was used for a reverse transcription reaction with iScript Reverse Transcription Supermix for RT-qPCR (Bio-Rad). One microliter product was used for real-time PCR with SsoAdvanced Universal Probes Supermix (Bio-Rad) and TaqMan primers/probe sets (Thermo Fisher Scientific) on a QuantStudio3 system (Thermo Fisher Scientific). Three different cDNA templates were tested in duplicate (5, 7).

**Whole transcriptome sequencing.** Whole transcriptome sequencing (RNA-Seq) of FReP cells was performed by the UCLA Technology Center for Genomics & Bioinformatics. Briefly, Illumina (HiSeq 2500, Illumina) 1 × 50 bp short reads were aligned to the transcriptome
Figure 8. Intratesticular implantation of CDKN2B-KD FReP cells in Fox Chase SCID Beige mice results in teratoma formation. (A) Gross appearance and histological evaluation (H&E staining) of adult Fox Chase SCID Beige mouse testes with intratesticular implantation of 1 × 10^6 cells were assessed 4 months after implantation. All implanted mouse testes are shown in Supplemental Figure 8 (n = 10 mice per group). (B) Teratoma formation in CDKN2B-KD FReP cell-implanted animals was evaluated by a pathologist and confirmed by immunofluorescent staining; teratoma formation from retrovirus-mediated BJ-iPSC-implanted animals (Figure 4A) was used as a positive control. Three germ layers are clearly identified by ectoderm – pigmented cells; mesoderm cartilage with type II collagen staining; and endoderm gland with the definitive endoderm marker forkhead box A2 (FOXA2) staining. Scale bars: 5 mm (black), 1 mm (blue), or 50 μm (orange).

derived from the University of California, Santa Cruz, Human Genome version 19 (hg19) (90). The results were submitted to the Gene Expression Omnibus (GEO GSE104830). Differentially expressed genes (fold change ≥±2) between FReP cells and BJ-derived iPSCs that were generated without genome integration (GEO SRR500985/6/7/8, ref. 23; and SRR1983694/5, ref. 24) were identified by the TopHat-Cufflinks package (25) through the Galaxy platform (26) using the default settings. DAVID Bioinformatics Resource 6.8 was used for functional annotation (27, 91). The differentially expressed genes that aligned with human proto-oncogenes and tumor suppressor genes, listed in the UniProt database (28), were displayed graphically by the package pheatmap (version 1.0.8) in R (version 3.4.1) (92).

**Western blot.** Cells were lysed using Pierce RIPA buffer supplemented with Halt Protease and Phosphatase Inhibitor Cocktail (Thermo Fisher Scientific). Western blotting was performed with antibodies against CDKN2B (catalog ab53034, Abcam) and GAPDH (product 5174S, clone D16H11, Cell Signaling Technology) to confirm the efficiency of CDKN2B KD at the protein level. Pierce ECL Western Blotting Substrate (Thermo Fisher Scientific) was used for development.

**Intramuscular implantation animal model.** Two-month-old male SCID mice (Charles River Laboratories) were anesthetized with inhaled 2% isoflurane and oxygen (1.5 L/min) and were maintained in an oxygen flow environment upon awakening. A 0.5-cm incision was recognized as a statistically significant difference (95). 0.05 (*) was considered a suggestive difference, while 0.01 (**), 0.001 (***), and 0.0001 (****) were used for nonparametric data. For all data presented in this article, P < 0.05 (*) was considered a suggestive difference, while P < 0.005 (**) was recognized as a statistically significant difference (95).

**Study approval.** All animal surgeries were performed under institutionally approved protocols provided by Chancellor’s Animal Research Committee at UCLA (protocol 2012-119).

**Data and materials availability.** All data generated or analyzed during this study are included in this article (and its supplementary information files). The RNA-Seq data of FReP cells were submitted to the NIH Gene Expression Omnibus (GEO GSE104830).

**Author contributions**

ZZ conceived and designed the study. ZZ, CL, PY, WJ, XP, and ZM performed the in vitro and in vivo experiments, data interpretation, and statistical analysis, except for the following: PH, GXC, and JKK performed the animal surgeries. XZ performed the pathological analysis of the teratomas. ZZ and CL wrote and revised the manuscript. EAB, CSH, and EC edited and proofread the manuscript. ZZ, KT, and CS supervised the entire study and all authors reviewed the final manuscript. The authorship order of the co-first authors was determined by flipping a coin.

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