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Combined modulation of polycomb and trithorax genes rejuvenates β cell replication

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Inadequate functional β cell mass underlies both type 1 and type 2 diabetes. β Cell growth and regeneration also decrease with age through mechanisms that are not fully understood. Age-dependent loss of enhancer of zeste homolog 2 (EZH2) prevents adult β cell replication through derepression of the gene encoding cyclindependent kinase inhibitor 2a (*INK4a*). We investigated whether replenishing EZH2 could reverse the agedependent increase of *Ink4a* transcription. We generated an inducible pancreatic β cell–specific *Ezb2* transgenic mouse model and showed that transgene expression of *Ezb2* was sufficient to increase β cell replication and regeneration in young adult mice. In mice older than 8 months, induction of *Ezb2* was unable to repress *Ink4a*. Older mice had an enrichment of a trithorax group (TrxG) protein complex at the *Ink4a* locus. Knockdown of TrxG complex components, in conjunction with expression of *Ezb2*, resulted in *Ink4a* repression and increased replication of β cells in aged mice. These results indicate that combined modulation of polycomb group proteins, such as EZH2, along with TrxG proteins to repress *Ink4a* can rejuvenate the replication capacity of aged β cells. This study provides potential therapeutic targets for expansion of adult β cell mass.

Introduction

Patients with type 1 or type 2 diabetes have inadequate functional β cell mass, a clinical need framing worldwide interest in the mechanisms controlling β cell growth and regeneration (1, 2). β Cell mass expands in response to increased metabolic demands associated with physiological growth, pregnancy, obesity, and insulin resistance in mice and humans (3-6), and increasing evidence suggests that this expansion is driven by replication of preexisting β cells (7–10). However, prior studies suggest that the capacity of β cells to replicate is age dependent, suggesting that restoring functional β mass in diabetic patients may require the manipulation of mechanisms that naturally limit the regenerative capacity of aging β cells (3, 11–13). Several studies have shown that increased transcription from the cyclin-dependent kinase inhibitor 2a (Ink4a) locus, which encodes the protein p16^{Ink4a}, an inhibitor of CDK4 activity (hereafter referred to as Ink4a), curtails β cell replication during aging (14, 15). Further evidence for a role for Ink4a-mediated control of β cell expansion was provided by genome-wide association studies linking the Ink4a/Arf locus to insulin insufficiency and risk of type 2 diabetes in humans (16, 17). Thus, identifying and controlling the mechanisms that regulate transcription of Ink4a locus during aging are the focus of intensive efforts and could be useful for promoting β cell regeneration.

Prior studies from our laboratories have demonstrated that members of the polycomb group (PcG) of proteins are involved in regulating transcription from the *Ink4a* locus during aging (18). PcG proteins exist in distinct polycomb repressive complexes that function sequentially to repress expression of target genes. Polycomb repressive complex 1 (PRC1) contains Bmi/ubiquitin ligase-Ring1B proteins, and polycomb repressive complex 2 (PRC2)

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contains the histone methyltransferase called enhancer of zeste homolog 2 (EZH2; refs. 18-20). PRC2 catalyzes the trimethylation of lysine 27 in the tail of histone H3 (H3K27me3), signaling the recruitment of PRC1, which then mediates the ubiquitination of lysine K119 in histone H2 (H2AK119), resulting in the repression of the Ink4a locus (21, 22). ChIP analysis quantified association of PcG proteins with the *Ink4a* locus in pancreatic islet β cells and showed high levels of PcG protein enrichment in islets from young mice that were significantly higher than those in islets from aged mice. The age-dependent loss of PRC1 and PRC2 at the islet Ink4a locus was accompanied by declining levels of polycomb proteins in β cells (18, 19). Moreover, the Ink4a locus showed increased enrichment of H3K4me3 and increased binding of the trithorax group (TrxG) protein complex that contains Mll1, a histone methyltransferase that catalyzes trimethylation of H3K4 (23-26). Thus, chromatin changes at the Ink4a locus result in age-dependent increases of *Ink4a* expression to attenuate adult β cell replication.

Inactivation of a conditional *Ezh2* allele in β cells leads to premature expression of *Ink4a* and severe reduction of β cell replication, demonstrating a crucial in vivo role for *Ezh2* in repressing *Ink4a* in islets (18). Natural reductions of *Ezh2* mRNA and protein levels in islets with aging could therefore account for derepression of the *Ink4a* locus in aged islets (19). Here, we tested whether replenishing *Ezh2* in islets during aging could prevent *Ink4a* derepression and reverse age-dependent declines in β cell replication. If so, expression of *Ezh2* in aged islets could rejuvenate the replicative capacity of β cells to promote regenerative expansion following β cell injury in adult mice.

Here, we report the generation and analysis of transgenic mice permitting conditional expression of *Ezh2* in adult pancreatic β cells. Induction of *Ezh2* in young adult mice was sufficient to repress *Ink4a* and stimulate β cell replication and regeneration. However, *Ezh2* induction in β cells of aged mice failed to repress

Ink4*a* and rejuvenate the capacity for replication. We show that this resistance to EZH2 results from enrichment of the Mll1-containing TrxG complex at the Ink4*a* locus. Combined knockdown of *Mll1* and activation of *Ezh2* was sufficient to repress Ink4*a* and increase β cell replication in aged mice. The principal elements of the PcG/TrxG/Ink4a pathway are conserved in human β cells, indicating that reprogramming of the Ink4*a* locus by modulating PcG-TrxG could be developed for use in diabetic patients.

Results

Inducible β cell-specific Ezh2 expression in vivo promotes β cell replication. Prior studies have shown that Ezh2 mRNA and protein levels in islet β cells decline with advancing age in mice and humans (19, 27). In aging mouse islets, ChIP revealed reduced association of EZH2 and histone H3K27me3 levels at the Ink4a locus, accompanied by increased levels of H3K4me3 and Ink4a mRNA expression (19, 28). In islets from humans with advancing age, we observed similar changes (Supplemental Figure 1, A-C; supplemental material available online with this article; doi:10.1172/ JCI69468DS1). Thus, evolutionarily conserved mechanisms linked to EZH2 reduction may constrain β cell replication with age. To test the possibility that conditional Ezh2 activation might be sufficient to rejuvenate islet β cell proliferation, we constructed a transgenic mouse line permitting conditional Ezh2 reexpression in aged mouse β cells. First, we created mice harboring a bidirectional tetracycline-responsive element (TRE) that controlled expression of a transgene encoding myc-tagged Ezh2 (Myc-Ezh2TG, referred to herein as EzTG) and β -gal (Figure 1, A and B; see Methods). Intercross of EzTG mice with a mouse line expressing reverse tetracycline transactivator (rtTA) in $\boldsymbol{\beta}$ cells directed from rat insulin promoter elements (*RIP-rtTA*; refs. 18, 19) generated bitransgenic progeny (RIP-rtTA; EzTG, referred to herein as bEzTG). Exposure of bEzTG mice to doxycycline (Dox) for 1 week induced expression of *EzTG* mRNA and protein (Figure 1, C and J) and β -gal (Figure 1D) in pancreatic β cells. By contrast, bEzTG mice without Dox exposure (Figure 1, D and F, and Figure 2A) or single RIP-rtTA or EzTG transgenic mice exposed to Dox did not show increased EzTG expression levels or β -gal (data not shown). Thus, construction of the bEzTG mice permitted studies of conditional *Ezh2* induction in islet β cells.

To investigate whether *Ezh2* reexpression was sufficient to stimulate adult β cell replication, we exposed 2-month-old bEzTG mice to Dox, when endogenous β cell EZH2 levels are nearly undetectable (29). In bEzTG mice exposed to Dox for 1 week (Figure 1E), we observed a 2-fold increase of the proliferation antigen Ki67 in islet insulin⁺ cells by immunostaining compared with that in littermate controls without Dox exposure (Figure 1, F and G). Exposure to Dox for 2 weeks (see Methods) led to a 50% increase in β cell mass, as assessed by morphometry (Figure 1H). Immunohistochemistry revealed a striking reduction of Ink4a in islet β cells from Dox-exposed bEzTG mice compared with controls (Figure 1I), a change confirmed by Western blotting (Figure 1J). Collectively, these data suggest that reexpression of *Ezh2* in young adult β cells is sufficient to reduce Ink4a and promote β cell replication.

Inducible β cell–specific Ezh2 expression in vivo is insufficient to induce replication of β cells in aged mice. Diabetes incidence increases with aging (19, 28), but the basis for this observation is unclear. Thus, we studied whether EzTG induction could lead to a similar β cell phenotype of increased replications in mice at more advanced age. Prior studies suggest that, by 8 months of age, islets lose



EzTG recruitment to the Ink4a locus in islets is age dependent. Based on resistance of islet β cell replication to *Ezh2* induction in aged bEzTG mice, we postulated that EZH2 association at loci such as Ink4a might be restricted by age. To test this possibility, we performed ChIP to assess the Ink4a locus in 2-month-old and 8-month-old bEzTG mice, respectively. As expected, ChIP revealed increased EZH2 association at the Ink4a locus in islets from Doxexposed 2-month-old bEzTG mice compared with islets from control bEzTG mice not exposed to Dox (Figure 3A; appropriate ChIP antibody and primer controls and validation are presented in Supplemental Figure 2A). Increased histone H3K27me3 levels at the Ink4a locus accompanied this EZH2 increase (Figure 3, B and C). Consistent with prior studies, we also measured increased association of the PRC2 complex protein EED, the PRC1 protein Bmi-1, and ubiquitinated H2A at the Ink4a locus in islets from Dox-exposed 2-month-old bEzTG mice (Supplemental Figure 2, B-D). By contrast, we failed to detect increased EZH2, EED, Bmi-1, or H3K27me3 at the Ink4a locus in islets from Dox-exposed 8-month-old bEzTG mice (Figure 3, G-K, and Supplemental Figure 2, E–G), despite increased *Ezh2* transgene expression (Figure 2E). Thus, the Ink4a locus is resistant to EZH2 association and Ezh2-dependent chromatin changes in aged islets.

We postulated that increased association of H3K4 methyltransferase Mll1 with the *Ink4a* locus could antagonize association of EZH2. ChIP analysis revealed relatively low levels of Mll1 and H3K4me3 association at the *Ink4a* locus in islets from 2-monthold mice, and treatment of bEZTG mice with Dox did not alter this association (Figure 3, D and E). Eight-month-old control and bEzTG mice had increased association of Mll1 and H3K4me3 at the *Ink4a* locus compared with 2-month-old mice (Figure 3,



Figure 1

Conditional expression of EZH2 in 2-month-old bEzTG mice promotes β cell replication through repression of Ink4a. (**A**) Schematics of the targeting cassette containing LacZ-TRE-EzTG. (**B**) Southern blotting to detect EzTG integration within the genome (lane 2–7), as compared with that in the WT control (lane 1). EzTG vectors were used as a positive control (lane 8–10). Expression level of (**C**) *EzTG* mRNA and (**D**) β -gal, as shown by LacZ staining in bEzTG islets treated with Dox for 1 week but not in littermate controls without Dox exposure (original magnification, ×20). (**E**) Dox feeding schemes: bEzTG mice were treated for 1 week with Dox food and drinking water, before the pancreata or the islets were harvested for experiments. For measuring β cell mass, 2- and/or 8-month-old (2M/8M) bEzTG mice were treated with an extended Dox procedure, with Dox treatment on week 1 and 3. Pancreatic tissue were harvest after 1 month. (**F** and **I**) Pancreatic sections from the control and Dox-treated 2-month-old bEzTG mice were immunostained with antibodies to insulin (green), (**F**) Ki67 (red), (**I**) Ink4a (red), and DAPI (blue) (original magnification, ×20). (**G**) Quantification of the percentages of Ki67-positive β cells with or without Dox exposure (5 pancreatic sections per animal). (**H**) β Cell mass (mg) measurement. (**J**) Western blotting and quantification of EzTG (Myc Ab) and Ink4a levels in isolated islets from indicated samples. *n* = 5–6 animals per group. **P* < 0.005.

I and J). These observations support the hypothesis that increased Mll1 and/or H3K4me3 levels at the *Ink4a* locus locus in aged islets could impair association of EZH2 with that locus and prevent repression of *Ink4a* by transgenic EZH2.

Mll1 occupancy at the Ink4a locus restricts Ezh2 recruitment in aged islets. We postulated that disrupting Mll1 association with the Ink4a locus might permit EZH2 recruitment to that locus in aged β cells. To test this possibility, we used siRNA to knockdown Mll1 levels in cultured islets from 8-month-old bEzTG mice (see Methods). To enhance assessment of β cells in cultured islets, we used nuclear Pdx1 immunostaining to quantify proliferating β cells that incorporated BrdU. Following exposure of islets to Dox, quantification of BrdU⁺Pdx1⁺ cells revealed that EZH2 induction was insufficient to increase β cell replication, similar to the observations in vivo (Figure 4, A and B). Knockdown of *Mll1* alone without *Ezh2* induction led to a modest reduction of Ink4a and increase in replication, but the combination of *Ezh2* induction and *Mll1* siRNA led to a 2.5-fold increase in BrdU incorporation (Figure 4, A and B). Consistent with these data, qRT-PCR and Western blotting studies demonstrated that *Ink4a* mRNA and protein levels were clearly reduced by the combination of *Ezh2* induction and *Mll1* siRNA in 8-month-old bEzTG islets (Figure 4,





Figure 2

Activation of EZH2 expression using 8-month-old bEzTG mice did not show any changes in β cell replication or regeneration. (A and D) Pancreatic sections from the control and Dox-treated 8-month-old bEzTG mice were immunostained with antibodies to insulin (green), (A) Ki67 (red), (D) Ink4a (red), and DAPI (blue) (original magnification, ×20). (B) Quantification of the percentages of Ki67-positive β cells with or without Dox exposure (5 pancreatic sections per animal). (C) β cell mass (mg) measurement. (E) Western blotting and quantification of EzTG (Myc Ab) and Ink4a levels in isolated islets. (F) STZ and Dox treatment scheme: Dox was administrated after single dose STZ (90 mg/kg) injection; tissues were harvested 7 days after injection. Immunostaining of insulin (green), Ki67 (red), and DAPI (blue) in pancreatic sections from bEzTG animals 1 week after STZ injection at (G) 2 months and (I) 8 month (original magnification, ×20). Percentages of Ki67-positive β cells with or without Dox exposure 1 week after mock or STZ treatment are quantified in (H) 2-month-old and (J) 8-monthold groups. n = 3-6 animals per group. *P < 0.05, ***P < 0.005.

C and D). In these Dox-exposed bEzTG islets, ChIP analysis revealed that simultaneous *Ezh2* induction and *Mll1* knockdown led to increased EZH2 association and H3K27m3 levels at the *Ink4a* locus locus, accompanied by reduced Mll1 and H3K4me3 levels (Figure 4, E–H). Together, these data support our view that a complex containing Mll1 protein restricts association of EZH2 with *Ink4a* in islets from aged mice.

Targeting TrxG complex by JmjD3 knockdown enhances Ezh2-induced β cell replication in aged islets. MLL proteins interact with histone H3K27 demethylases (19), including UTX and JmjD3 (31, 32). JmjD3 associates with established PcG targets, including Ink4a, and regulates H3K27me3 levels (33). However, it remains unclear whether Mll1 associates with JmjD3 in β cells. We observed a modest increase in H3K27me3 at the Ink4a locus and reduced Ink4a mRNA levels in 8-month-old bEzTG islets with Mll1 knockdown (Figure 4, C and F), consistent with the possibility that reduction of TrxG proteins might affect H3K27 demethylases and Ink4a expression in aged islets. To identify proteins associated with Mll1 in the TrxG complex, we performed Mll1 immunoprecipitation in mouse insulinoma Min6 cells. Mll1 immunoprecipitation revealed its association with JmjD3, along with another core component of the TrxG complex, RbBP5 (Supplemental Figure 3, A and B). Since the recruitment of Mll1 at the Ink4a locus increases with age, we postulated that, as an interaction partner of Mll1, the association of JmjD3 with the Ink4a locus might follow a similar pattern. We found that the levels of JMJD3 mRNA increased with age in human and mouse islets (Figure 5A and data not shown). Consistent with this finding, ChIP analysis with JmjD3 antibody revealed increased association of JmjD3 with the Ink4a locus of human and mouse islets, as a function of age (Figure 5B and Supplemental Figure 3, C and D). Thus, increased JmjD3 demethylase association accompanies reduced H3K27me3 levels at the Ink4a locus in aging islets. To



Figure 3

EZH2 transgene is recruited to Ink4a locus in 2-month-old, but not 8-month-old, bEzTG islets. (A) Schematic representation of the Ink4a locus, with blue regions marked 1–4 indicating the amplified regions in the ChIP studies. Representative ChIP analysis for the indicated antibodies at the Ink4a locus in islets isolated from (B-F) 2-month-old and (G-K) 8-month-old groups.

test the possibility that JMJD3 might regulate induced EZH2 activity in islets, we used siRNA targeting to knockdown JMJD3 in aged bEzTG islets. Following exposure of 8-month-old bEzTG islets to Dox (resulting in *Ezh2* induction) and to an siRNA targeting *JMJD3*, immunostaining of Ki67 and Pdx1 demonstrated a 5-fold increase in β cell replication (Figure 5, C and D, and Supplemental Figure 3D). This effect on β cell replication is similar to that observed following simultaneous knockdown of Mll1 and Ezh2 induction in these islets. Similar to Mll1 knockdown experiments, ChIP analysis revealed that simultaneous Ezh2 induction and JMJD3 knockdown led to increased EZH2 association and H3K27m3 levels at the Ink4a locus, accompanied by reduced Mll1 and H3K4me3 levels (Figure 5, E-J). However, a combined knockdown of *Mll1* and *JmjD3* in islets from 2-month-old mice did not show any difference in proliferation, indicating that this mechanism is age dependent (Supplemental Figure 3, E and F). Together, our findings suggest a 2-step mechanism of Ink4a-mediated aging that includes attenuation of PcG protein levels and accumulation of TrxG complexes, including JmjD3, at the *Ink4a* chromatin in aged β cells.

Dominant function of TrxG at the Ink4a locus is a general mechanism in aging cells. We postulated that the regulation of Ink4a by PcG and TrxG proteins might be a general mechanism mediating Ink4a-based replicative senescence. To test this, we studied mouse embryonic fibroblasts (MEFs); like islet β cells, MEFs accumulate Ink4a with passaging (34, 35). In serially passaged C57BL/6 MEFs, levels of EZH2 declined, whereas Ink4a mRNA and protein levels increased, changes accompanied by reduced proliferation, as quantified by Ki67 immunodetection (Supplemental Figure 4, A and B). Like those in pancreatic β cells, EZH2 and H3K27me3 levels at the Ink4a locus decreased in late-passage MEFs, while Mll1 and H3K4me3 levels increased, confirming prior studies (Supplemental Figure 4, C–G, and ref. 27). To test whether TrxG restricts

EZH2-dependent Ink4a repression in late-passage MEFs, we transfected these MEFs with DNA vectors permitting myc-EZH2 misexpression, Mll1 siRNA-mediated knockdown, or both. Consistent with the results in pancreatic β cells, we observed that simultaneous Mll1 knockdown and Ezh2 induction increased fibroblast proliferation, accompanied by a 40% reduction of *Ink4a* mRNA levels. (Figure 6). Mll1 knockdown alone slightly enhanced BrdU incorporation, while Ezh2 induction alone did not detectably alter BrdU labeling (Figure 6B). Therefore, rejuvenation of the replicative capacity of β cells to promote regenerative expansion may require a 2-step approach, including dissociation of TrxG proteins and replenishment of the PcG proteins at the Ink4a chromatin. Taken together, these data support our findings and conclusions from studies on pancreatic β cells and suggest that combined activity of TrxG and PcG to regulate Ink4a transcription could be a general approach to promote cell regeneration.

Discussion

To achieve functional β cell restoration in diabetic patients, the mechanisms regulating β cell replication have been intensively investigated. Using transgenic mice to offset the loss of endogenous EZH2, we have advanced the concept of regenerative plasticity in adult β cells. In young adulthood, activation of EZH2 expression in β cells was sufficient to repress *Ink4a* and increase the replication of β cells. This is consistent with previous observations that the β cell mass in young adults can expand to adapt to metabolic demands accompanying pregnancy and obesity (27). Thus, results here and in prior work suggest that, even in the absence of ongoing replication, β cells in young adults retain the capacity for replication and expansion. By contrast, aged mice are resistant to adaptive changes in β cell mass, and replenishing EZH2 alone in aged mice was not sufficient to repress the *Ink4a* locus. Our find-



Figure 4

Knockdown of MII1 facilitates EZH2 recruitment to the Ink4a locus and increased β cell replication. (**A**) 8-month-old bEzTG islets were treated with scrambled siRNA (100 pM), Dox (1 µg/ml), *MII1* siRNA (100 pM), and both Dox and *MII1* siRNA for 48 hours. Islets sections were immunostained for Ki67 (red) and Pdx1 (green) (original magnification, ×20). (**B**) BrdU was added in culture 48 hours before harvest to measure proliferation. Percentage of BrdU-positive β cells in each experiment group (5 pancreatic sections per animal). (**C**) Real-time qRT-PCR of *Ink4a, Ezh2*, and *MII1* mRNA in each group. (**D**) Western blotting and quantification of EzTG (detected by Myc Ab), Ink4a, and β -tublin protein levels. (**E**–I) Representative ChIP result for the indicated antibodies at the Ink4a locus in islets treated as in **A**. Data represents 3 experiments. **P* < 0.05.

ing that chromatin changes at the *Ink4a* locus accurately reflect β cell replicative capacity validates investigations of *Ink4a* expression to promote β cell regeneration and, more generally, justifies studies of age-dependent chromatin changes in islet β cells.

Our findings also suggest that efforts to regenerate β cells will be advanced by identifying the signaling pathways that regulate expression and activity of PcG and TrxG factors in aging β cells. Previously, we showed that PDGF signaling regulates β cell EZH2 levels and is an important conserved mechanism in sustaining β cell replication during neonatal expansion of mouse and human islets (13). Here, we observed increased recruitment of the Mll1-JmjD3 protein complex to the *Ink4a* locus, coinciding with reduced levels of PDGF signaling and EZH2 decline, accompanied by increased levels of JmjD3 in the β cells. Moreover, findings with bEzTG mice here suggest that activation of *Ink4a* expression by the Mll1-JmjD3 complex may prevent EZH2-dependent repression of









Figure 6

Knockdown of MII1 combined with EZH2 expression results in increased proliferation of later passage MEFs. (A) Passage 13 MEFs were transfected with scrambled siRNA (100 pM), pcDNA Myc-EZH2 (2 µg), MII1 siRNA (100 pM), and both pcDNA Myc-EZH2 and MII1 siRNA for 48 hours. BrdU was added in culture 48 hours before immunostaining for BrdU (red) and DAPI (blue) (original magnification, ×20). (B) Percentage of BrdU-positive MEFs in each experiment group. (C) Real-time qRT-PCR of Ink4a, Ezh2, and *Mll1* mRNA in each group. n = 3experiments. *P < 0.05, **P < 0.01.

the Ink4a locus in aged β cells. Further studies are needed to determine whether PDGF signaling or other pathways regulates the age-dependent changes in Mll1 and JmjD3 that limit β cell replication.

The loss of H3K27me3-repressive marks at the Ink4a locus could not be solely attributed to the decline in the binding of EZH2, and our study highlights an active mechanism of removal of H3K27me3 marks involving JmjD3. We show that JmjD3 and Mll1 act together to modify 2 distinct lysines: Jmjd3 demethylated H3K27, while Mll1 methylated H3K4, resulting in transcriptional activation of the Ink4a locus. This mechanism of H3K27me3 removal appears to be a very tightly regulated process, highlighted by the fact that, while the levels of H3K27 methyltransferase EZH2 decline with aging, there is a concomitant increase in the levels of H3K27 demethylase, JmjD3, in β cells. The importance of JmjD3 is underscored by studies in MEFs, which show that ectopic expression of JmjD3 leads to increased expression of Ink4a and induction of replicative senescence (28). Our experiments indicate that loss of JmjD3 in β cells from aged animals leads to repression of *Ink4a* and induction of β cell replication, suggesting a critical role for active removal of H3K27me3 marks in the regulation of Ink4a locus.

Previous studies have demonstrated that *Ink4a* repression by PcG proteins is a well-conserved mechanism across cell types (35) as well as during iPS cell generation (36). Our experiments with late-passage MEFs show that reexpression of EZH2 was insufficient to repress *Ink4a* unless coupled with loss of the Mll1 complex, in a manner similar to pancreatic β cells. Thus, TrxG-PcG mechanisms governing replicative senescence elucidated here may be generally relevant to creating regenerative strategies for multiple types of tissues and cells.

Methods

pBi-G-Myc-Ezb2 construct. pBI-G tet vector is used to express EZH2, which contains a bidirectional tet-responsive promoter containing TRE between 2 minimal CMV promoters that control the expression of β -gal and EZH2. The plasmid was cloned as follows. A mouse EZH2 full-length cDNA fragment was obtained by BssHII digestion and blunt-ended with T4 DNA polymerase, followed by a NotI digestion from the parental plasmid from an ATCC clone (Image clone ID: 6817366). This sequence was inserted into pcDNA3-Myc vector in frame with Myc-tag at N-terminal, with the vector

treated with EcoRI digestion, blunt-ended with T4 DNA polymerase, and then digested with NotI. pcDNA3-Myc vector was generated by insertion of the Myc tag into into KpnI/BamHI sites. The myc-EZH2 sequence was then isolated from pcDNA3-MycEZH2 by KpnI digestion, blunt-ended with T4 DNA polymerase, digested with XhoI and then cloned into pBi-g Tet vector with the vector treated with PstI digestion, blunt-ended with T4 DNA polymerase, and followed with a SalI digestion.

Generating inducible bEzTG mice. A 9-Kb DNA fragment that contains LacZ-TRE-MycEZH2 sequences was then obtained by AseI digestion of pBiG-MycEZH2 and injected into F2 donor embryos of C57BL/6 × DBA/2 and FVB/N animals to generate transgenic animals. Founder mice with the correct insert were then crossed into Rip-rtTA mice to generate inducible bEzTG mice with bEzTG double transgene.

Mouse husbandry and genotyping. Animals were fed ad libitum and kept under a 12-hour-light/dark cycle. DNA extracted from the tails was used for PCR-based genotyping using standard methods. Primers used for genotyping bEzTG were as follows: forward 5'-CACGCTGTTTTGACCTC-CATAG-3' and reverse 5'-TGGAAAATCCAAGTCACTGGTG-3'.

Dox administration. Dox was administrated both in drinking water (2 g/l) supplemented with sucrose (0.5%) shielded from light in red bottles and food pellets (200 mg/kg, Bio-Serv). Dox food and water were changed every 2 to 3 days. Gender-matched littermates with bEzTG genotype on normal diet and drinking water were used as controls. For proliferation index analysis, pancreata were harvested after 2 weeks of Dox treatment. For β cell mass analysis, pancreata were harvested after 1 month of feeding, with Dox treatment throughout week 1 and 3 and normal chow and water throughout week 2 and 4 (Figure 2A).

Regeneration studies. A single dose of 90 mg/kg body weight STZ (Sigma-Aldrich) was prepared fresh in 0.1 M citrate buffer (pH 4.5) and injected intraperitoneally into animals in different age groups. STZ-treated animals were exposed to Dox or control diet/water immediately after the injection for 1 week before the pancreata were harvested for proliferation index analysis.

Southern blotting. Genomic DNA from control and transgenic mice was digested with PvuII, transferred to nylon membrane, and incubated with Myc-EZH2 full-length probe. pcDNA-Myc-EZH2 was digested with KpnI/Xhol, and 2.6-kb length Myc-EZH2 sequence was purified from the gel and used as a template to generate the probe. Probe preparation and detection were performed using the DIG High Primer DNA Labeling and Detection Starter Kit (Roche) according to manufacture's instructions.

LacZ staining. LacZ staining of isolated islets was performed in islets from bEzTG and littermate control mice 1 week after Dox exposure. In brief, freshly isolated islets were fixed at 4°C in 4% paraformaldehyde, prepared in 100 mM PBS for 10 minutes, followed by 3 washes of cold PBS with 0.1% Triton X-100 and 2-hour incubation in PBS solution containing 5 mM potassium ferricyanide, 5 mM potassium ferrocyanide, 2 mM MgCl₂, and 1 mg/ml X-gal at 4°C. After staining, images were collected with a dissecting microscope equipped with a Zeiss camera.

RNA isolation, RT-PCR, and real-time qPCR. RNA from islets was isolated with the RNeasy Mini/Micro Kit (Qiagen) according to the manufacturer's instructions. 200 ng–1 µg RNA was used for preparation of cDNA using SuperScript III Reverse Transcriptase (Invitrogen) with the oligo dT primers. The cDNA was then analyzed with regular PCR or real-time quantitative PCR (qPCR). Real-time qPCR was performed using ABI7900HT (Applied Biosystems), with initial denaturation at 95 °C for 20 seconds, followed by 50 cycles of 94 °C for 1 second and 60 °C for 20 seconds, which was then followed by a dissociation stage. The expression levels of each transcript were normalized to the housekeeping gene cyclophilin A (*PPIA*). Each real-time PCR experiment shown is a representative from at least 3 independent experiments; for each experiment, islets were pooled from 3 to 4 mice per specified group.

For lists of RT-PCR and real-time qRT-PCR primers used for mRNA expression analysis, see Supplemental Tables 1 and 2.

Immunoprecipitation and Western blotting. 2.5 µg anti-Mll1 antibody (05-765, Millipore) or control IgG was used for each immunoprecipitation. Antibodies were immobilized to protein G Sepharose (Millipore) for 2 to 3 hours and washed with cold nondenaturing buffer (20 mM Tris-HCl [pH 8], 137 mM NaCl, 10% glycerol, 1% NP-0, 2 mM EDTA). The antibody-coated beads were then incubated with Min6 cell extracts prepared in nondenaturing buffer overnight at 4°C on a rotator. Beads were washed 3 times in ice-cold nondenaturing buffer and eluted in 1X SDS-PAGE gel-loading buffer. The immunoprecipitates were analyzed by Western blotting with antibodies against JmjD3 (07-1434, Millipore), RbBP5 (A300-109A, Bethyl Laboratories), and control Dnmt3a (IMG-268A, Imgenex).

For Western blotting, isolated islets were lysed in cell dissociation buffer (Invitrogen) supplemented with protease inhibitor cocktail (Calbiochem). Protein concentration was measured with Dc protein assay (Bio-Rad), and protein samples (or immunoprecipitates) (10 μ g-30 μ g) were loaded to each lane of the SDS-PAGE gel as previously described (18, 19, 27, 37). Target protein level was quantified against the housekeeping protein β -tublin using ImageJ. Each Western blot shown is a representative from 3 independent experiments using different islet preparations.

Immunohistochemistry. Immunohistochemistry was performed as previously described (8). Briefly, pancreatic tissue was fixed in 4% formaldehyde and processed for paraffin embedding. 5-µm sections were deparaffinized and rehydrated and then subjected to antigen unmasking and permeabilization. The slides were then blocked with 3% BSA in TBST (0.1% Tween 20) and incubated with primary antibody overnight at 4°C. Primary antibodies used for study included guinea pig anti-insulin (1:400; Dako), mouse anti-Ki-67 (1:40; BD Biosciences), mouse anti-BrdU (1:2; RPN-202, GE Healthcare/Amersham), mouse anti-p16 (diluted 1:250; sc-1661, Santa Cruz Biotechnology Inc.). The slides were then washed and incubated with donkey- and goat-derived secondary antibodies conjugated to FITC or Cy3 (1:200; Jackson ImmunoResearch Laboratories). Slides were mounted with Vectashield with DAPI (Vector Laboratories) prior to imaging using a Leica DM6000 microscope and Openlab software (Improvision). The immunofluorescence data presented are representatives of at least 5 animals per group in each group.

For analysis of the β cell mass and proliferation index, 3–6 mice were analyzed in each group. For β cell mass, pancreatic sections were stained

for insulin and scanned using a Leica DM6000 microscope. Montages of the pancreatic area by DAPI staining and β cell area by insulin staining were made with ImageJ. The cross-sectional areas of pancreata and β cells were determined using an automated program on Openlab software. Proliferation index was calculated using Ki67 (or BrdU) staining as a marker for proliferation and insulin (or Pdx1) staining for β cells.

ChIP protocol. ChIP was performed as previously described (38). In brief, isolated or cultured islets (150-200 islets per ChIP) were treated with 1% paraformaldehyde at room temperature for 10 minutes for crosslinking. The islets were washed and suspended in ChIP lysis buffer with protease inhibitors and sonicated with Bioruptor (Diagenode). The chromatin was then precleared with protein A/G beads. Meanwhile, 10 µl protein A or G beads were incubated with 2 to 5 mg of the antibodies anti-H3K27me3 (Diagenode), anti-Ub-H2A (05-678, Millipore; the antibody is an IgM and therefore requires a bridging antibody [12-488, Millipore]), anti-H3K4me3 (Diagenode), anti-Bmi-1 (05-637, Millipore), anti-EZH2 (clone AC22, no. 3147, Cell Signaling), anti-Mll1 (05-765, Millipore), anti-JmjD3 (07-1534, Millipore), or normal mouse IgG as a control at 4°C for 2 hours on a rotator. The antibody-bound bead mix was then washed and incubated with precleared chromatin overnight at 4°C on a rotator. After immunoprecipitation, the chromatin was eluted from the beads and subjected to reverse crosslinking, and the DNA was purified. The ChIPed DNA was quantified as percentage bound/input with real-time PCR, performed using the ABI Master Mix Kit (Life Tech) and the ABI 7900 HT Fast Real-Time PCR Machine (Life Tech). Each ChIP experiment was performed 3-5 times, using independent pooled islet isolations from 4 to 8 mice. 21. The real-time PCR primers used to amplify the Ink4a locus in the ChIP DNA spanned throughout the locus and were labeled 1-4, as indicated in Figure 3A.

Mouse islet isolation and culture. Islets were isolated using the Liberase/ DNase Enzyme Mix (Roche Diagnostics) as described previously (39). In brief, the pancreas was inflated with the enzyme mix through the common bile duct and digested at 37° C. Islets were enriched by gradient and hand picked. 100–200 islets were cultured in Easy-Grip Tissue Culture Dishes (35 mm × 10 mm, Falcon) in DMEM (Cellgro) supplemented with 10% FBS. Islet cells were transfected after overnight resting in culture using Lipofectamine 2000 (Invitrogen) according to the manufacturer's instructions. The islets were harvested 48 hours after transfection and processed for immunostaining, protein extraction, RNA extraction, and ChIP analysis. For proliferation analyses in islets, cultured islets were incubated with BrdU (1 µg/ml) and/or Dox (2 µg/ml) after transfection 48 hours before they were harvested. The islets were then fixed in 4% paraformaldehyde, embedded in HistoGel (Thermo Scientific), and processed for paraffin-embedded sections.

Human islet studies. Human islet samples were obtained from healthy, nondiabetic organ donors, deceased due to acute traumatic or anoxic death and obtained from the National Diseases Resource Interchange or Integrated Islet Distribution Program as previously described (8). Equilibrated human islets from different age groups, ranging from 8 months to 70 years, were processed for ChIP and real-time qRT-PCR analysis similar to the mouse islets. ChIP results in humans were normalized to GAPDH locus. For *JMJD3* mRNA and ChIP results (Figure 5, C and D), samples were grouped into juvenile (<10 years) and adult (\geq 10 years). ChIP results for EZH2 and histones are shown for each age group. qRT-PCR results were normalized to *Actb.* For *Ezb2* mRNA, samples were grouped into young (<20 years) and old (\geq 20 years).

MEF culture. MEFs were isolated from E13.5 C57BL/6 embryos as previously described (28). In brief, embryos were harvested in cold sterile PBS, and embryo head and organs (everything inside body cavity) were carefully removed. The remainder tissue was minced and trypsinized to make single cell suspension. Trypsin was quenched by adding MEF medium (DMEM

with 10% FBS). The cells at P0 were centrifuged and cultured in dishes coated with gelatin. Early-passage MEFs were passaged every 2 to 3 days using trypsin; however, the proliferation slowed down over a few passages and could be passed every 4 to 5 days or weekly toward the later passages.

Statistics. All data are expressed as mean \pm SEM. Mean and SEM values were calculated from triplicates (at least) of representative experiments. Statistical significance was determined by an unpaired, 2-tailed Student's *t* test. A *P* value of less than 0.05 indicated statistical significance.

Study approval. All animal experiments were approved by the Animal Research Committee of the Office for the Protection of Research Subjects at UCLA.

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