Notch signaling dynamically regulates adult β cell proliferation and maturity

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**Graphical abstract**

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Notch signaling is an intercellular signal transduction pathway that regulates cell fate decisions. It has been studied extensively during embryonic development, where it controls cell proliferation and differentiation. However, the role of Notch signaling in the mature pancreas has been less explored.

**Introduction**

β Cell mass and function adapt to the insulin requirements of the organism to maintain euglycemia across a wide range of pathophysiology, such as insulin resistance, pregnancy, or aging (1). Molecular mechanisms that enable β cell adaptation to these stressors are not yet fully understood; insufficient functional adaptation, including β cell death and dedifferentiation, becomes clinically apparent with the onset of type 2 diabetes (T2D) (2, 3). With the continued increase in obesity, novel therapeutically tractable pathways that regulate β cell adaptation are sought to reverse β cell dysfunction in T2D.

Notch mediates cell-fate decisions via juxtacrine interactions between neighboring cells expressing Notch receptors (Notch1–4) and ligands (Jagged1/2, Dll1/3/4). Mice lacking the common transcriptional effector of all Notch signaling, Rbpj, or other Notch pathway components are embryonically lethal, but even prior to lethality, Notch mutant mice showed abnormal pancreata (4, 5). In fact, both Notch gain and loss of function impaired β cell differentiation (6, 7), underscoring the necessity of Notch signaling for normal development, but the potential role of Notch in the adult endocrine pancreas remained mostly unexplored, which was surprising with the subsequent identification of NOTCH2 as a T2D susceptibility locus (8).

We have recently shown that liver Notch activity is increased in diet-induced obesity (DIO) mouse models and in T2D patients (9) and contributes to excess hepatic glucose production and fatty acid synthesis (10, 11). These data led us to hypothesize that Notch may be similarly reactivated in the stressed β cell, consistent with other “developmental” pathways in the dedifferentiated β cell (3). Here, we find that Notch signaling is present at low levels in fully developed β cells, but increased in islets cultured in hyperglycemic conditions or isolated from obese mice. Persistent β cell Notch signaling appears detrimental to function, as forced Notch activation impaired glucose-stimulated insulin secretion (GSIS) in isolated mouse or human islets and glucose intolerance in β cell–specific Notch gain-of-function mouse models. Conversely, we observed improved glucose tolerance with genetic inhibition of β cell Notch action. Mechanistically, we found that Notch interfered with MafA-Kat2b association, which induced MafA proteasomal degradation (9) and contributed to excess hepatic glucose production and fatty acid synthesis (10, 11). These data led us to hypothesize that Notch may be similarly reactivated in the stressed β cell, consistent with other “developmental” pathways in the dedifferentiated β cell (3).

**Results**

Notch signaling is dynamically regulated in developed pancreatic β cells. As a first step to evaluating a potential postdevelopment role of β cell Notch signaling, we determined the absolute expression levels of Notch receptors/ligands in islets isolated from WT adult mice (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI98098DS1). This analysis revealed relatively high expression levels of Notch receptors/ligands in islets, consistent with prevalent Rbpj staining in β cells (Figure 1A), suggesting potential for ongoing β cell Notch signaling. To assess Notch activation in the adult pancreas, we used transgenic Notch reporter (TNR) mice that express GFP under the control of a Rbpj consensus sequence (12). Chow-fed TNR mice showed readily detectable Notch activity in a subset of β cells (Figure 1B), but trivial staining in other islet endocrine cells (Supplemental Figure 1B). We also observed an enrichment of β cell transcripts (i.e., Ins2) in FACS-sorted, Notch-active GFP+ islet cells (Supplemental Figure 1C). We next used another Notch reporter mouse model, with a more persistent fluorescent reporter (Histone2b-Venus) (13), to confirm these data — integrating both data sets, we found the islet Notch activity was essentially confined to β cells (Supplemental Figure 1D), but at generally lower
levels than Notch activity detected in endothelial or duct cells (Supplemental Figure 1E), consistent with earlier observations (14).

We next evaluated whether β cell Notch activity was regulated by metabolic stimuli. We used low-dose streptozotocin (STZ) treatment to render TNR mice hyperglycemic, which increased β cell Notch activity (Figure 1C and Supplemental Figure 1F). This was confirmed by increased expression of the canonical Notch target Hes1 in the surviving β cell population (Figure 1D). We attributed increased Notch activity to hyperglycemia, as opposed to an STZ-induced injury response, as high-glucose exposure also resulted in increased GFP protein levels and Hes1 expression in isolated TNR islets (Figure 1, E and F), consistent with increased Notch signaling in islets from hyperglycemic NOD mice (15). Similarly, we found higher expression of Hes1 and other Notch transcriptional targets in islets isolated from DIO mice, as compared with Chow-fed mice (Figure 1G), which corresponded with increased β cell Hes1 staining (Figure 1H). Thus, we conclude that Notch signaling is active in adult β cells and enhanced in conditions associated with increased insulin demand (obesity, partial ablation of β cell mass, and hyperglycemia).

β cell Notch activation impairs GSIS, leading to glucose intolerance. To test whether Notch may contribute to β cell dysfunction, we generated β cell–specific Notch gain-of-function (β-NICD) mice, which express the constitutively active Notch intracellular domain (NICD), by crossing Rosa26-NICD (7) with RIP-Cre (16) transgenic lines. Despite unchanged body weight (Supplemental Figure 2A), Chow-fed β-NICD mice were mildly glucose intolerant as compared with Cre− controls (Figure 2A). These differences were exacerbated by aging (Figure 2B) or high-fat diet (HFD) feeding (Figure 2C), but completely reversed with simultaneous Rbpj deletion (Figure 2D), suggesting glucose intolerance was mediated by canonical Notch signaling. β-NICD mice showed unchanged fasting insulin levels.
To exclude the impact of Notch activation during the early postnatal period, we generated tamoxifen-dependent (TAM-dependent), β cell–specific (17) Notch gain-of-function (MIP-β-NICD) mice. MIP-β-NICD mice were indistinguishable from controls before TAM treatment, but consistent with data from β-NICD mice, showed increased glucose excursion after TAM treatment that was exacerbated by HFD feeding (Supplemental Figure 2C). In sum, these data suggest that forced Notch activation in fully developed β cells is sufficient to impair functional response to obesity-induced insulin resistance.

Notch activity leads to a specific loss of β cell MafA in mouse and human islets. To determine how Notch affects β cell function, we took a candidate approach, focusing initially on key transcriptional regulators of GSIS, such as Pdx1, MafA, and Nkx6.1. Of these, we found a profound reduction in MafA staining in pancreatic sections, islets, and primary β cells from β-NICD– and TAM-treated MIP-β-NICD mice (Figure 3, A and B, and Supplemental Figure 3, A and B), but importantly, not in β-NICD:Rbpj mice (Supplemental Figure 3C). Notch-mediated loss of β cell MafA appeared relatively specific, without effects on Pdx1 or Nkx6.1 (Figure 3C, Supplemental Figure 3D and not shown), and also did not lead to a compensatory response in MafB, which localizes in glucagon+ cells equally in control and β-NICD mice (Supplemental Figure 3, E and F). Consistent with the hypothesis that decreased MafA is the mechanism of impaired GSIS with β cell Notch activation, shRNA-mediated Mafa silencing failed to exacerbate Notch inhibition of GSIS (Supplemental Figure 3G). Finally, NICD transduction in human islets also resulted in loss of Mafa (Figure 3D) and impaired GSIS (Figure 3E), as well as lower Pdx1, but without changes in MafB and Nkx6.1 (Supplemental Figure 4, A–C).

Notch causes MafA degradation by preventing its association with Kat2b. Intriguingly, Notch activation reduced MafA protein without affecting Mafa expression (Supplemental Figure 5, A and B), suggesting a posttranscriptional effect. To test this hypothesis, we generated a stable MIN6 line to allow doxycycline-dependent expression of a Notch1 gain-of-function mutant (N1ΔE), which does not require ligand binding, but still requires γ-secretase processing to render NICD (18). As hypothesized, N1ΔE led to loss of MafA (Figure 3F), but not in the presence of a γ-secretase inhibitor (GSI) (Supplemental Figure 5, C and D) or in cells that express dominant-negative mastermind-like 1 (DNMAML), which inhibits its function of the MAML transcriptional coactivator essential for Rbpj-dependent Notch signaling (Supplemental Figure 5E). Using this model, we confirmed that acute Notch activation rapidly decreases MafA protein (Figure 3G) without change in Mafa mRNA (Figure 3H).

We next observed that MG132-mediated proteasome inhibition preserved MafA protein levels (Figure 4A), which led us to hypothesize that Notch increases GSK3β activity, known to trigger MafA proteasomal degradation (19, 20). N1ΔE, however, did not affect GSK3β activation (Supplemental Figure 5F), although GSK3β inhibition prevented Notch-mediated MafA ubiquitination (Supplemental Figure 5G) and degradation (Supplemental Figure 5H). Similarly, whereas exogenous MafA was also degraded with Notch activation, the GSK3β-resistant Mafa-S65A mutant (21) was protected from Notch-induced degradation (Supplemental Figure 5I). These results suggest that Notch-induced proteasomal degradation of MafA is downstream of GSK3β-mediated phosphorylation.

The histone acetyltransferase Kat2b has been reported as protecting MafA from proteasomal degradation in cancer cells (20); as Kat2b can also interact with NICD (22), we hypothesized that Notch may affect MafA-Kat2b interaction, leading to MafA degradation. Indeed, Notch activation prevented normal Kat2b–MafA association (Figure 4B), and Kat2b overexpression in MIN6 cells protected MafA from Notch-mediated degradation (Figure 4C). Conversely, Kat2b silencing in MIN6 cells induced loss of MafA (Figure 4, D and E) without changing Mafa expression (Supplemental Figure 5J) and recapitulated the impaired GSIS associated with Notch activation (Figure 4F). In sum, these data show that Notch disrupts MafA-Kat2b interaction, which induces MafA proteasomal degradation and impaired β cell function (Figure 4G).

Notch activation and reduced maturity in proliferating β cells. Profiling experiments have reported reduced gene expression of maturity markers in proliferating β cells (23). In line with this idea, we observed lower MafA protein in BrdU+ cells from WT pregnant females (Figure 5A and Supplemental Figure 6A), but normal Pdx1, Neurod1, or Ucn3 (Figure 5B). We found a similar effect in MIN6 cells, which lose MafA when undergoing chromatin condensation prior to cell division, but retain Pdx1 and Nkx6.1 (Supplemental Figure 6B). Consistent with a Notch-MafA axis to regulate β cell maturity and proliferation, we observed increased Notch-active β cells in pregnant females as compared with virgin females (Figure 5C) as well as in the highly proliferative β cells remaining after STZ treatment (Figure 1, C and D, and Supplemental Figure 6C). In fact, the population of KI67+ proliferating β cells in pregnant TNR mice was enriched in GFP+ cells (Figure 5D); consistently, MafA levels were modestly reduced in GFP+ cells (Figure 5E).

These results suggest that a transient increase in Notch activity accompanies loss of β cell maturity in the course of normal β cell self-replication. We next asked whether persistent Notch activation causes loss of β cell maturity — indeed, islets from β-NICD mice showed a transcriptional signature of immaturity (decreased Mafa, Glp1r, and Ucn3 and increased Ldha; Supplemental Figure 6, D and E). Similar data were observed in TAM-treated MIP-β-NICD mice (Supplemental Figure 6F). Reduced Mafa and maturity in Notch-active β cells could arise from a feed-forward loop of Maf transcription factors (24), as previously shown for Mafa (25), or because chronic metabolic abnormalities in Notch gain-of-function mice repressed Mafa. To distinguish these possibilities, we treated islets derived from MIP-β-NICD mice with chronic 4-hydroxytamoxifen (4-OHT) treatment, which also revealed reduced Mafa (Supplemental Figure 6, G and H). In sum, these data indicate that β cell Notch activation destabilizes MafA, which in turn represses the β cell maturation program, but does not result in complete loss of cell identity, as lineage-tracing experiments did
not reveal β cell transdifferentiation or dedifferentiation in β-NICD mice (Supplemental Figure 6I).

Notch induces β cell proliferation, leading to increased β cell mass. The positive correlation between Notch activity and β cell proliferation in TNR mice led us to test causation. We observed an increase in β cell mass in β-NICD mice, due to a shift toward larger islet size (Figure 6A and Supplemental Figure 7A). We attributed increased β cell mass in β-NICD mice to increased β cell proliferation (Figure 6, B and C), detectable as early as P14 (Figure 6D), as β cell death was unchanged (Supplemental Figure 7B). β Cell mass remained higher in β-NICD mice even at 1 year of age (Figure 6E), when β cell proliferation rates were low and not different from those of control mice (Supplemental Figure 7C), suggesting that Notch-induced β cell mass acquisition may be maintained through adulthood.

To determine the necessity of intact Notch signaling for proliferation, we generated β cell–specific Notch loss-of-function not reveal β cell transdifferentiation or dedifferentiation in β-NICD mice (Supplemental Figure 6I).

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Figure 3. Notch activation leads to loss of MafA in mouse and human β cells. (A) Representative images of pancreatic sections from β-NICD and Cre control mice, stained with antibodies directed against insulin and MafA (n = 6 mice/group, quantification shown in Supplemental Figure 3C). (B) Western blots in islets isolated from β-NICD and Cre control mice. (C) Representative images of immunofluorescence of fixed MIN6-rTTA3 cells transfected with doxycycline-inducible NICD-HA (or GFP control) after stimulation with doxycycline for 24 hours, with quantitation of percentage of NICD+ cells with detectable MafA, Pdx1, or Nkx6.1 (n = 3 biologic replicates). Boxed region in second row is shown amplified; arrowheads indicate Notch+ cells with absent MafA. (D) MafA and insulin staining in dispersed human β cells transduced with lentivirus expressing GFP or NICD. Representative images from 3 individual donors selected out of a total sample size of 6 donors. (E) GSIS in islets from healthy human donors transduced with lentivirus expressing GFP or NICD. Results are expressed as fold increase in insulin secretion (16.8 mM glucose/2.8 mM glucose) and reflect data from 5 individual donors. **P < 0.01, ratio paired t test. (F) Western blots from MIN6-rTTA3 cells with stable integration of doxycycline-dependent N1ΔE-myc (or luciferase control) after stimulation with doxycycline for 24 hours. Doxy, doxycycline. (G) Western blots from MIN6-rTTA3 cells with stable integration of doxycycline-dependent N1ΔE-myc after stimulation with doxycycline for the indicated amount of time. (H) Mafa expression in MIN6-rTTA3 cells with stable integration of doxycycline-dependent N1ΔE-myc (or luciferase control) after stimulation with doxycycline for 24 hours (n = 3 biologic replicates). Scale bars: 20 μm. All data are shown with group means.
Figure 4. Notch causes MafA degradation by preventing its association with Kat2b. (A) Representative immunofluorescence images of fixed MIN6 cells transiently transfected with NICD-HA, with or without MG132 (n = 4 independent experiments). (B) Western blots from immunoprecipitates derived from 293T cells with stable integration of the rTTA3 transcriptional transactivator, transfected with MafA-myc, Kat2b-Flag, and Tet-On NICD-HA, with or without stimulation with doxycycline for 4 hours. (C) Representative images of immunofluorescence of fixed MIN6-rTTA3 cotransfected with Tet-On NICD-HA and GFP or Kat2b-Flag after stimulation with doxycycline for 8 hours, and quantitation of percentages of NICD+GFP+ or NICD+Kat2b+ cells with detectable MafA (n = 3 independent experiments, ~250 cells analyzed per sample). *P < 0.05, 2-tailed t test. (D) Western blots from MIN6 cells transduced with lentivirus encoding shRNA targeting Kat2b (#1 and #2) or scrambled control (Scr). (E) Immunofluorescence images of fixed MIN6 cells transduced with lentivirus encoding shRNA targeting Kat2b or scrambled shRNA (Scr). Representative images from 2 independent experiments. (F) GSIS from MIN6 cells transduced with lentivirus encoding shRNA against Kat2b or scrambled control in medium containing low (1 mM, LG) or high (25 mM, HG) glucose. Results from a representative experiment with 6 replicates per condition. Experiment was performed 3 times. ***P < 0.001, 1-way ANOVA and Dunnett’s multiple comparisons post hoc test. (G) Model of Notch regulation of MafA-Kat2b interaction and MafA proteasomal degradation. Scale bars: 20 μm. All data are shown with group means.
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The β-DNMAML mice, which showed decreased β cell proliferation at P14 (Figure 6F). To establish whether Notch regulation of β cell proliferation was cell autonomous, we isolated islets from MIP-β-NICD and MIP-β-DNMAML mice, then treated them with 4-OHT to induce recombination. Consistent with in vivo results, forced Notch activation increased glucose-induced proliferation, while genetic blockade of β cell Notch activation reduced proliferation (Figure 6G and Supplemental Figure 7D). Importantly, we observed unchanged proliferation capacity when cells were cultured in low glucose, where proliferation rates remained low (Supplemental Figure 7E). Similarly, Notch gain of function enhanced proliferation during pregnancy (Figure 6H), whereas pregnancy-induced β cell proliferation was diminished in Notch loss-of-function mice (Figure 6I). We conclude from these results that endogenous Notch activation augments β cell proliferation in times of need, but simultaneously suggest that Notch is neither necessary nor sufficient to force proliferation-incompetent (i.e., aging, low-glucose growth conditions) β cells to divide.

β Cell Notch inhibition improves obesity-induced glucose intolerance. We next circled back to our original observations and

Figure 5. Notch activation and reduced maturity in proliferating β cells. (A) Representative images of pancreatic sections from day 15 (D15) pregnant (D15) WT mice and quantitation expressed as average MafA fluorescence intensity of BrdU+ and BrdU− populations (n = 5 mice/group). Individual cells are plotted in Supplemental Figure 6A. (B) Representative images of pancreatic sections from pregnant (D15) mice stained for BrdU as well as antibodies directed against insulin and Pdx1, Neurod1, or Ucn3 (n = 5 mice/group). (C) Representative images of pancreatic sections from virgin and pregnant (D15) TNR mice, with quantitation of percentage of GFP+ β cells (n = 4–5 mice/group). (D) Representative images of pancreatic sections from pregnant (D15) TNR mice showing Ki67, insulin, and GFP reporter staining (top), with increased magnification of the boxed region (bottom). Arrows indicate Ki67+ β cells. Quantitation of percentage of GFP+ β cells or percentage of GFP+Ki67+ β cells (n = 5 mice/group; overall count of 3995 total β cells and 188 Ki67+ β cells). (E) Quantitation of average MafA fluorescence intensity in GFP− and GFP+ β cells from chow-fed TNR mice, normalized to average value in GFP− cells (n = 5 mice). Approximately 1000 cells quantified per pancreas. Scale bars: 20 μm. All data are shown with group means. *P < 0.05; **P < 0.01; ***P < 0.001, 2-tailed t test.
hypothesized that blocking Notch signaling in β cells may increase MafA stability and improve obesity-induced glucose intolerance. Indeed, expression of DNMAML reversed loss of MafA seen in primary β cells challenged with high glucose and palmitate (Figure 7A). Consistently, DNMAML increased the half-life of exogenous MafA in cycloheximide-treated MIN6 cells (Figure 7B).

To determine whether β cell–specific Notch loss of function could reverse loss of MafA in HFD-fed mice (Supplemental Figure 6–hypothesized that blocking Notch signaling in β cells may increase MafA stability and improve obesity-induced glucose intolerance. Indeed, expression of DNMAML reversed loss of MafA seen in primary β cells challenged with high glucose and palmitate (Figure 7A). Consistently, DNMAML increased the half-life of exogenous MafA in cycloheximide-treated MIN6 cells (Figure 7B).

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feeding was initiated at 24 weeks of life (Supplemental Figure 8, D–G), in contrast with MIP-CreERT mice fed HFD when β cells were more proliferative (26). With this experimental paradigm, MIP-β-DNMAML mice demonstrated improved glucose tolerance (Figure 7C). As predicted, islets from these mice showed lower Notch activity (Supplemental Figure 8H), but also indications of greater β cell maturity (Figure 7, D and E).

To ensure reproducibility, we created a second, inducible Notch loss-of-function mouse model, MIP-β-Rbpj mice, which...
showed normal glucose tolerance when fed a chow diet (Supplemental Figure 8I), but improved glucose tolerance as compared with Cre− controls when challenged with HFD feeding (Figure 7F). Consistent with data from MIP-β-DNAML mice, islets from MIP-β-Rbpj mice had decreased Notch activation (Supplemental Figure 8J), with a parallel increase in MafA levels, leading to a prematurity β cell expression profile (Figure 7, G and H). Finally, as predicted, given the delay in HFD-induced insulin resistance until β cells were less proliferative, both Notch loss-of-function models showed β cell mass and β cell proliferation that were the same as those of Cre− controls and therefore had identical fasted serum insulin levels (Supplemental Figure 8, I–K, and not shown). These data indicate that Notch inhibition stabilizes MafA, leading to increased β cell maturity and improved glucose tolerance in the face of obesity-induced metabolic challenge.

Discussion

Reactivation of “embryonic” signaling pathways can occur in environmentally stressed β cells (3). For instance, increased Notch signaling has been shown in hypoxia (28) or when islets are compromised in the setting of pericyte ablation (29). Since these same stressors also lead to β cell dysfunction, cause and effect cannot be disentangled, but here we show that unrestrained Notch activation is detrimental to β cell maturity and leads to disrupted GSIS in isolated mouse and human islets as well as impaired glucose tolerance in vivo associated with increased systemic demand for insulin. Reciprocally, genetic inhibition of Notch activation maintains β cell function when mice are challenged by HFD feeding independently of Notch’s well-established role in pancreatic endocrine development (6, 7, 30).

Our data also indicate that constitutive Notch activation cannot induce dedifferentiation, unlike forced activation of Hedgehog signaling (31), nor is Notch activation sufficient to trigger β cell replication, but rather, it leads to a higher proliferative rate under conditions associated with β cell division. These data suggest that Notch activation pushes stressed β cells further away from the status quo, but whether the positive effect of Notch signaling on β cell proliferation is a direct consequence of Notch activation, a byproduct of loss of β cell maturity, or both is not yet known. These findings do, however, support the idea of β cell function being opposed to β cell proliferation (32) and also support a model in which the proliferating population of β cells is both Notch active and functionally immature, with a readout of destabilized MafA. These data are consistent with in vitro work showing Notch activation in proliferating human β cells (33, 34) as well as studies on proliferating β cells in vivo (23), which showed repression of the Pdx1/NeuroD/MafA network, but not gene expression of these canonical β cell transcription factors or loss of other maturity markers, such as Ucn3 (35). Of these, MafA has been consistently shown to be essential for β cell function and survival in the adult pancreas (36–38). Various circumstantial data suggest that partial loss of MafA activity may represent commitment to β cell division: (a) forced MafA expression in pancreatic progenitors increased expression of cell cycle inhibitors and blunted cell proliferation (39); (b) MafA only becomes essential for β cell function after weaning (36); and (c) MafA expression is low during the postnatal proliferative wave (40) and in hyperproliferative human insulinomas with altered MENIN expression (41). However, total ablation of MafA in whole body or pancreas-specific knockout models does not result in hyperproliferative β cells — on the contrary, loss of β cell mass was reported (36, 38). One possible explanation — that loss of MafA may lead to compensatory MafB expression, reported to mediate β cell proliferative adaptation to pregnancy (42, 43) and β cell maturation (44, 45) — did not hold true in Notch-active β cells. This may indicate that dynamic regulation of MafA activity is required for normal regulation of β cell maturity and proliferation, perhaps by Notch effects on Kat2b-MafA interaction. Thus, our data are broadly consistent with the protective role of MafA for β cell adaption to stress (46–48), but further studies are necessary to test MafA dependence on Notch’s various effects on β cell biology as well as to determine Kat2b interaction partners in β cells in various pathophysiologic states.

We observe heterogeneous Notch activation in β cells, consistent with differential proliferative capability of distinct β cell populations (49, 50); in fact, the proliferative “bottom β cell population” identified by Rui and colleagues shows increased Hes1 expression (50). Notch activation is at minimum a readout of this β cell heterogeneity, but likely accounts for variability in cell-to-cell MafA and resultant propensity to proliferate. Although we have not yet identified the source of the Notch signal, our preliminary data point to β cell-specific upregulation of specific Notch ligands of the Delta-like and Jagged families in response to hyperglycemia, which suggests the possibility of stressed β cells sending out a call for proliferation. But our data cannot exclude other endocrine or non-endocrine signal–sending cells; similarly, as our Notch loss-of-function mouse models focused on common downstream signaling components (MAML, Rbpj), future studies will be necessary to identify the specific Notch receptor activated in β cells.

In sum, we find active Notch signaling in the “normal” β cell, but with 2 disparate outcomes: in young mice, Notch activation represents an adaptive mechanism to increase β cell mass induced by insulin resistance by permissive action on proliferation, but in aged mice with lower proliferative capability (51) or with chronic insults, such as obesity, Notch activation locks β cells into an immature and dysfunctional state — a maladaptive compensatory response. Thus, Notch loss of function enhances β cell maturity and improves glucose tolerance. These data parallel findings in liver (9) and other insulin-sensitive tissues (52, 53), where similar Notch “reactivation” is causal to insulin resistance (10, 11, 54–56). As Notch is therapeutically accessible (57), with small molecules and biologics in the pipeline for cancer (58), these results provide the groundwork to repurpose these agents to address the dual pathologies — insulin resistance and deficient insulin secretion — that characterize T2D.

Methods

Animals. RIP-Cre: Tg(Ins2-cre)23Herr (16), MIP-CreERT: B6.Cg-Tg(Ins1-cre/ERT)1Lphi/J (17), R26-NICD: Gt(Rosa)26Sorcre;R26ltm1(LacZ)Jpg (7), R26-DN-MAML1: Gt(Rosa)26Sorcre;R26ltm1(MAML1);Exp1Forest (59), and Rbpjfl/fl; Rbpjnull (6) mouse lines were maintained on a C57BL/6J background. TNR: Tg(Cp-EGFP;2Sgaia/Reja) mice were maintained in homozygosity to avoid epigenetic silencing on a C57BL/6J and SJL/J mixed background (12). H2B-Venus notch reporter: Tg(Cp-HISHTH2B/Venus)47Hadj mice were maintained in homozygosity on a FVB background (13). Mice were
weaned and maintained on standard chow (Purina Mills, catalog 5053) or HFD (Harlan Laboratories, TD.06414) as stated. Adult mice (8–12 weeks old) were used for experiments, unless stated otherwise. No differences in body weight or glucose tolerance were found between WT and RIP-Cre or MIP-CreERT mice (not shown). Male mice were used for all experiments described, with the exception of experiments involving pregnancy, and ex vivo GSIS, where islets from mice and female mice were used without distinction.

**Antibodies and chemicals.** All antibodies used can be found in Supplemental Table 1. STZ, TAM, 4-OHT, doxycycline hyclate, MG132, cycloheximide, BrdU, and LiCl were obtained from MilliporeSigma. CHIR 99021 was obtained from Tocris Bioscience.

**STZ and TAM treatment.** STZ was dissolved in cold citrate buffer, 0.1M pH 4.5, and injected within 10 to 15 minutes of preparation, once daily at a dose of 50 mg STZ/kg body weight for 5 consecutive days. Mice were sacrificed 18 days after last STZ injection, with fed glucose levels of 450–520 mg/dl. TAM was dissolved in corn oil (MilliporeSigma) and sterile filtered; both Cre controls and CreERT mice were treated with 100 mg TAM/kg body weight once daily for 3 consecutive days.

**Islet isolation.** Islets were routinely isolated by collagenase P (Roche Applied Science) digestion of whole pancreas (except for electron microscopy experiments), followed by Histopaque density-gradient centrifugation, as described (60). When required, islets were maintained ex vivo in 5.5 mM glucose RPMI 1640 supplemented with 10% FBS and 1% penicillin-streptomycin. For islet dispersion experiments, isolated islets were incubated overnight and dissociated with Accutase (Corning). Dispersed cells were plated onto glass coverslips and cultured on islet medium.

**Human islet studies.** Human islets from 6 nondiabetic donors (ages 25–57 years old, BMI 23.6–33.5 kg/m²) were obtained through the Integrated Islet Distribution Program (IIDP) via NIH-approved centers. Islets were cultured in PIM(S) media supplemented with 5% human serum (Prodo Labs).

**Glucose tolerance tests and GSIS assays.** MIN6 cells were incubated in 1 mM glucose Krebs-Ringer-HEPES (KRH) buffer (129 mM NaCl, 5 mM NaHCO3, 4.8 mM KCL, 1.2 mM KH2PO4, 1.2 mM MgSO4, 2.5 mM CaCl2, 10 mM HEPES pH 7.4, 0.1% BSA) for 1 hour, then transferred to 20 mM glucose for 1 hour. For islet GSIS, after overnight culture in 5.5 mM glucose RPMI, groups of 6 similar size islets were incubated for 1 hour in 2.8 mM glucose KRH buffer, subjected to 2.8 mM and 16.8 mM glucose stimulation, and supernatants collected. For total insulin content, islets were transferred to high-salt buffer (2.15 M NaCl, 10 mM NaH2PO4, 40 mM Na2HPO4, 1.8 mM EDTA pH 7.4) and sonicated for 2 minutes in a Bioruptor sonicator (Diagenode). Insulin in supernatant was detected by ELISA (Merckodia). Total islet DNA content was determined by quantitative PCR (qPCR) using a standard curve of defined DNA values. Glucose tolerance tests (GTTs) were performed as previously described (10).

**Cell lines and lentivirus.** Culture of insulinoma-derived MIN6 cells and lentivirus generation with the use of 293T cells were performed as described (61). For generation of a MIN6 cell line expressing a transcranial transactivator (MIN6-rTTA3), cells were transduced with lentivirus obtained with the pLenti CMV rTA3 Blast vector (Addgene, catalog 26429) and a blasticidin (5 μg/ml) selection. Vectors for shRNA-mediated knockdown of Mafa or Kat2b were generated by cloning targeting sequences (Supplemental Table 2) into pLKO.1-TRC cloning vector (Addgene, catalog 10879). To generate stable MIN6-rTTA3 cells for doxycycline-mediated expression of N1ΔE-myc or luciferase, N1ΔE cDNA was cloned from pCS2 Notch1 ΔEMV-6MT (Addgene, catalog 41737) into pLVX-Tight-Puro vector (Clontech) with pLVX-Tight-Luc (Clontech) used as a negative control. Lentiviral particles generated from these vectors were transduced into MIN6-rTTA3 cells, which were selected with 1 μg/ml puromycin (MilliporeSigma). MIN6 cells stably expressing DNAML-GFP or GFP control were generated after lentivirus infection and FACS sorting. All vectors used in this study can be found in Supplemental Table 3.

**Lentivirus-mediated transduction.** Lentivirus was generated in 293T cells, as previously described (61). Titration was performed by qPCR. Intact or dispersed islets were transduced with 10 transducing units (TU)/cell, assuming that an average islet contains 1000 cells, in complete medium enriched with 8 μg/ml sequabrene (MilliporeSigma). Imaging and GSIS experiments in human islets were performed 96 hours after lentivirus infection.

**Western blotting and PCR.** Lysates were resolved by SDS-PAGE, followed by Western blot and visualization using the ECL Western Blotting Detection Kit (GE Healthcare Bio-Sciences). RNA was isolated with TRizol (Thermo Fisher), cDNA obtained with the High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher), and qPCR performed using Power SYBR Master Mix (Thermo Fisher). Primer sequences used are detailed in Supplemental Table 2.

**Immunofluorescence, β cell mass determination, BrdU, and TUNEL assay.** Tissue was fixed in 4% paraformaldehyde/PBS at 4°C for 2 to 4 hours, followed by overnight incubation in 30% sucrose/PBS. Tissue was embedded in OCT compound (Sakura) and flash frozen in dry ice-chilled isopentane. Tissue sections 5 μm thick were processed according to standard procedures. For MafA and Hes1 immunostaining, slides were incubated for 20 minutes at 70°C with HistoVT One (Nacalai) for antigen retrieval. For BrdU staining, mice were injected intraarteriole with 100 mg/kg BrdU (MilliporeSigma) 5 hours before sacrifice. After standard processing of the tissue, slides were incubated at 37°C in 2N HCl for 30 minutes, followed by incubation in 0.1 M sodium borate buffer pH 8.5 for 10 minutes, prior to incubation in primary antibodies, then donkey secondary antibodies (Jackson Immunoresearch), both diluted in PBS containing 5% normal donkey serum, 1% BSA, and 0.1% Triton X-100. TUNEL assay was performed with the In Situ Cell Death Detection Kit, Fluorescein (Roche Applied Sciences) following the instructions from the manufacturer, and pancreas sections further stained with insulin and DAPI. The β cell mass analyses and immunofluorescence in MIN6 cells dispersed or intact islets were performed as previously described (60).

**Confocal microscopy.** For confocal microscopy assays, Axio Observer Z1 with LSM 710 scanning module was used (Zeiss). Images were obtained with a ×40 Zeiss Plan-Apochromat oil objective. Imaging was performed in a single confocal microscopy session for each experiment. For each session, the photomultiplier voltage settings (below 600V) and laser transmission (≤ 2%) for each fluorophore were determined to maximize the dynamic range of the signal. Controls were used for each experiment to confirm specific signals. All images were obtained in a 1024 × 1024 pixel format. For microscope operation and image gathering, ZEN (Zeiss) software was used.

**Quantification of immunofluorescence experiments.** Quantitative analysis of fluorescence intensity for nuclear antigens (MafA, Pdx1, or Hes1) was performed with images gathered from pancreas tissue slides. For this, raw multichannel LSM files were processed with ImageJ (NIH), and the channel of interest subjected to background correction using the “rolling ball” algorithm implemented in ImageJ.
The region of interest (ROI) was then drawn on 20–30 random nuclei per image, while visualizing the insulin channel to select β cells. The signal intensity for the ROI of the channel of interest (nuclear staining) was obtained per image and then averaged per pancreas (5–10 images per pancreas). Results shown are the average of 3–6 mice per group. The average pixel intensity values were referenced to the values obtained for the control of each experiment and expressed as AU. For qualitative analysis (Ki67, BrdU, MafA expression in vitro), a threshold was set using the Threshold function of ImageJ, and positive and negative cells were counted. Images were processed with ZEN2; the only manipulation performed was linear contrast stretching to attain the optimal dynamic signal range for each set of images. Representative pictures shown for these experiments have quantified values approximated to the final average.

Transmission electron microscopy. Islets were isolated with 0.1 mg/ml Liberase TL (Roche) and handpicked in HBSS containing 0.2% BSA and 2.8 mM glucose. Isolated islets were immediately fixed at 4°C for 3 hours with 2% paraformaldehyde, 2.5% glutaraldehyde, and 3 μM CaCl₂ in 0.1M sodium cacodylate buffer pH 7.4. Samples were processed as described (62). For quantification analyses, the area occupied by insulin vesicles and the β cell cytoplasmic area were measured with ImageJ using 50–60 images per mouse.

Islet diameter. Freshly isolated islets were maintained in ice-cold RPMI with 2% normal calf serum, and images were gathered with an Olympus IX70 microscope. Individual islet area was determined using ImageJ, and results converted to diameter.

Statistics. Results are shown as mean ± SEM. Differences between 2 groups were calculated using a 2-sided Student’s t test, ratio paired Student’s t test, or Mann-Whitney U test as indicated. Differences between multiple groups and a control were calculated by 1-way ANOVA followed by Dunnett’s multiple comparisons post hoc test. A P value of less than 0.05 was considered statistically significant.

Study approval. The Columbia University Institutional Animal Care and Utilization Committee approved all animal procedures. Human islets were harvested from deceased donors without any identifying information. Informed consent and IRB approval were obtained at islet isolation centers.

Author contributions AB conducted experiments and contributed to the experimental design, discussion, and writing of the manuscript. CZ conducted experiments and contributed to the review and editing of the manuscript. LS contributed to the experimental design, discussion, and review and editing of the manuscript. UBP contributed to the experimental design, discussion, and writing of the manuscript.

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