Spatiotemporal patterns of multipotentiality in Ptf1a-expressing cells during pancreas organogenesis and injury-induced facultative restoration

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SUMMARY
Pancreatic multipotent progenitor cells (MPCs) produce acinar, endocrine and duct cells during organogenesis, but their existence and location in the mature organ remain contentious. We used inducible lineage-tracing from the MPC-instructive gene Ptf1a to define systematically in mice the switch of Ptf1a+ MPCs to unipotent proacinar competence during the secondary transition, their rapid decline during organogenesis, and absence from the mature organ. Between E11.5 and E15.5, we describe tip epithelium heterogeneity, suggesting that putative Ptf1a+Sox9+Hnf1β+ MPCs are intermingled with Ptf1a+Sox9LO proacinar progenitors. In the adult, pancreatic duct ligation (PDL) caused facultative reactivation of multipotency factors (Sox9 and Hnf1β) in Ptf1a+ acini, which undergo rapid reprogramming to duct cells and longer-term reprogramming to endocrine cells, including insulin+ β-cells that are mature by the criteria of producing Pdx1HI, Nkx6.1+ and MafA+. These Ptf1a lineage-derived endocrine/β-cells are likely formed via Ck19+/Hnf1β+/Sox9+ ductal and Ngn3+ endocrine progenitor intermediates. Acinar to endocrine/β-cell transdifferentiation was enhanced by combining PDL with pharmacological elimination of pre-existing β-cells. Thus, we show that acinar cells, without exogenously introduced factors, can regain aspects of embryonic multipotentiality under injury, and convert into mature β-cells.

KEY WORDS: Multipotent pancreas progenitors, Facultative progenitors, Acinar-to-endocrine transdifferentiation, Injury, Mouse, Lineage tracing

INTRODUCTION
The inherent plasticity potential of adult pancreas is especially interesting considering the possibility of controllably reprogramming various cell types into β-cells as an alternative to embryonic stem cell differentiation and transplantation therapy for diabetes. Pancreatic multipotent progenitor cells (MPCs) are the basis for generating a functional pancreas with appropriate cell allocations and structures. Understanding the ontogeny and intermediate states of MPCs could provide insight into whether the differentiated adult pancreatic cells could be challenged to return to an embryonic multipotency under stimuli such as physiological injury or chronic diabetes.

The early pancreatic epithelium contains MPCs producing several transcription factors (TFs), including Ptf1a, associated with commitment toward pancreatic fate, and controlling early organ growth (Pan and Wright, 2011). At the beginning of morphogenetic maturation [around embryonic day (E) 12], the epithelium becomes compartmentalized into ‘tip’ and ‘trunk’ domains (Zhou et al., 2007; Villasenor et al., 2010). Genetic lineage-tracing via Cpa1CreER suggested that MPCs with the rudimentary signature Pdx1+Ptf1a+Myc+Hpf1a+ are located at the tip domain, but rapidly disappear with the start of the secondary (2°) transition (Zhou et al., 2007), and the trunk epithelium between E12.5 and E15.5 was proposed to harbor bipotential progenitor pools for duct and endocrine cells (Zhou et al., 2007; Schaffer et al., 2010). Lineage-tracing analysis using Hnf1βCreER (Solar et al., 2009) and Sox9CreER (Kopp et al., 2011), however, suggested that trunk epithelium contains a progressively reducing number of MPCs, even after overt tip-trunk segregation. While the Hnf1β population is multipotent during the early 2° transition, but largely bipotential at later stages, relatively low numbers of Sox9+ cells were proposed to feed all pancreatic lineages throughout embryogenesis, suggesting persistence of Sox9+ MPCs after the 2° transition (Kopp et al., 2011). Because Sox9+ and Hnf1β+ cells were localized to the trunk epithelium, with Cpa1CreER cells at the tips, the precise location and number, and molecular signature, of MPCs during and after the 2° transition remains unclear.

The existence of adult facultative progenitors capable of regenerating endocrine cells under special stimuli has been contentious (Granger and Kushner, 2009). Recent lineage tracing in mice, however, yielded the surprising suggestion that many terminally differentiated cell types could be reprogrammed to other cell-types, either using exogenous factors or under injury conditions. Xu et al. (Xu et al., 2008) concluded that facultative Ngn3+ (Neurog3+ – Mouse Genome Informatics) endocrine progenitors, producing β-cells ex vivo, appeared in/at the duct following pancreatic duct ligation (PDL). By contrast, lineage-tracing results with duct/centroacinar-specific CreER lines (Hhf1βCreER, Hes1CreER, Sox9CreER) provided evidence against duct and centroacinar cells deriving endocrine cells after PDL (Solar et al., 2009; Kopinke et al., 2011; Furuyma et al., 2011; Kopp et al., 2011). Detailed studies are required to determine rigorously the origin of the Ngn3+ duct cells and if they progress to islet/β-cells in vivo.
Because of their vast number, acinar cells could be an attractive source for replenishing β-cells to offset losses from disease or injury. Acinar-to-endocrine conversion was suggested with primary acinar cells cultured in vitro when treated with growth factors (Baeyens et al., 2005; Minami et al., 2005). In vivo lineage-tracing using ElastaseCreERTM however, showed no evidence of such conversion upon regeneration/injury with partial pancreatectomy (Ppx), PDL, or cerulein-induced pancreatitis (Desai et al., 2007). Strikingly, overexpression of three β-cell TFs – Pdx1, Ngn3, MafA – apparently turned acini into β-like cells in vivo, suggesting convertibility under strong stimuli (Zhou et al., 2008). It remains unclear whether all acinar cells are equally susceptible to such conversion, and if it could be activated without genetic intervention or under injury conditions that might exist in diabetes or pancreatitis.

Ptf1a is expressed in early bud MPCs, with an instructive role in distinguishing pancreatic fate from the adjacent organs (Kawaguchi et al., 2002). In the 2° transition pancreatic epithelium, Ptf1a production is dynamically regulated. Over time, its activity changes from driving an MPC program to directing tip cells into a proacinar state. Moving from MPC to proacinar behavior is proposed to be linked to the switching of Ptf1a co-regulatory proteins in the trimeric PTF1 complex, from PTF1RBP-J to PTF1RBP.JL (Masui et al., 2007). An outstanding issue is whether small numbers of Ptf1a+ MPCs persist during/after the 2° transition, or if adult Ptf1a+ acini could somehow re-engage (aspects of) an embryonic Ptf1a-driven MPC program, to adopt facultative progenitor activity.

Here, we report a knock-in tamoxifen-inducible Ptf1aCreERTM line for the dissection of gene regulatory networks critical for maintaining the MPC state and controlling tip-trunk compartmentalization. We reasoned that lineage-tracing analysis performed using the MPC-instructive factor Ptf1a would provide more certainty in addressing the intrinsic potentiality (mono, bi, or multi) of pancreatic progenitor populations during organogenesis. We describe a dynamic heterogeneity within the tip epithelium, based on distinct TF signatures, which changes during tip-trunk segregation concordant with the shift from multipotency to unipotency in Ptf1a+expressing cells. Pulse-labeling-type lineage-tracing using Ptf1aCreERTM showed that the number of Ptf1a+ MPCs decreased progressively and rapidly during organogenesis, with rare MPCs present late into the 2° transition, and perinatal/adult stage Ptf1a+ cells produce only acinar cells. However, Ptf1a+ acinar cells were activated by PDL to become facultative progenitors, express several embryonic multipotency factors and become competent to produce duct and endocrine cells, including mature β-cells. Acinar-to-endocrine conversion was detected in substantial numbers only 2 months post-PDL, but their formation was enhanced by eliminating pre-existing β-cells with a selective toxin, streptozotocin (STZ). This is the first study to demonstrate, with in vivo lineage-tracing, that acinar cells give rise to endocrine cells under injury-induced reprogramming paradigms and without additional transcription factors or signaling molecules.

MATERIALS AND METHODS

Mice

Ptf1aCreERTM was generated using recombinase-mediated cassette exchange (Long et al., 2004) (supplementary material Fig. S1A). R26RlacZ and R26R-EYFP mice were described previously (Soriano, 1999; Sirinivas et al., 2001). Animals and embryos were PCR-genotyped. Experiments were under protocols approved by Vanderbilt University IACUC.

Tamoxifen administration and injury procedures

Tamoxifen (Tam, Sigma T-5648) was 30 mg/ml in corn oil (Sigma C8267). Tamoxifen administration and dosage used were described in detail in results section. PDL was performed as described (Xu et al., 2008). Streptozotocin (STZ; 10 mg/ml in sodium citrate pH 4.5) was given intraperitoneally (100 mg/kg) for 2 consecutive days at post-PDL D3.

Tissue preparation and immunostaining

Embryonic or adult pancreas was fixed (2-4 hours, 4% paraformaldehyde, 4°C), and for cryosections, washed twice in cold PBS, sucrose-equilibrated (30%, 4°C, overnight), and embedded in OCT compound (Tissue-Tek, Sakura). Paraflin sections, whole-mount β-galactosidase detection, H&E staining, processing was as described (Kawaguchi et al., 2002). Immunofluorescence used 10 μm cryosections (Kawaguchi et al., 2002). Antibodies used are listed in supplementary material Table S1.

Data collection, morphometric and statistical analysis

Images from Zeiss confocal (LSM 510 META upright) or Apotome were analyzed with LSM Image Browser and Zeiss Axiovision 4.8 software, respectively. Systematic quantitation analysis was performed by Photoshop CS3 and NIH ImageJ software.

Short-term lineage tracing, E13.5 and E14.5: labeling of the acini, ducts or endocrine cells was manually scored as percentage of each EYFP+ cell-type of total cell number in each compartment, using all field of view, every other section (20 μm apart; ~50% total pancreas). E18: every sixth section (60 μm apart; ~16% total pancreas) was counted. Juvenile/adult mice: 150-200 random fields from 40 evenly spaced sections (100 μm apart; ~10% of total pancreas) were analyzed. For E18.5, and juvenile/adult pancreata, percentage labeling was calculated based on EYFP+ area over total area for each cell type. Head/tail pieces of PDL pancreata were separately embedded and analyzed. All fields of view from every 100 μm section (~40-45; ~10% total pancreas) ensured coverage of all regions. At least three to four pancreata per stage/group were analyzed. Two-tailed Student’s t-test used a significance of P<0.05.

RESULTS

Generation and characterization of Ptf1aCreERTM mice

Given that Ptf1a is expressed in early pancreatic MPCs (Kawaguchi et al., 2002), and later at E12 is seemingly restricted to MPC/proacinar progenitors in the tip epithelium of the remodeling epithelial plexus (Zhou et al., 2007), it was critical to determine quantitatively the dynamics of this shift from multipotential to unipotential behavior. To lineage-trace Ptf1a+expressing cells at different stages of pancreas organogenesis, we derived Ptf1aCreERTM knock-in mice expressing CreERTM under endogenous Ptf1a promoter/enhancer elements (supplementary material Fig. S1A). CreERTM production recapitulates endogenous Ptf1a expression, with nuclear translocation induced in Ptf1a+ and Cpa1+ acinar cells within 24 hours of tamoxifen (Tam) administration at E15.5 (supplementary material Fig. S2A,B).

Testing Ptf1aCreERTM mice with reporter alleles (Rosa26lacZ [R26RlacZ] or Rosa26EYFP [R26REYFP]) and Tam administered at designated developmental stages allowed indelible marking of the progeny of Ptf1a+ cells throughout subsequent development. Recombination of R26RlacZ was strictly tamoxifen-dependent and no pancreatic β-gal+ cells were detected in non-CreERTM mice (supplementary material Fig. S3A,B; Fig. 1C,C¢). EYFP+ labeling, while single 3 mg Tam doses at E16.5 and E18.5 resulted in ~50-60% labeling of acinar cells (supplementary material Fig. S3D; Fig. 3B-D), lower labeling (~10-40%) occurred with single 3 mg Tam doses at E16.5 and E18.5 resulted in ~50-60% labeling of acinar cells (supplementary material Fig. 3B-D). Lower labeling (~10-40%) occurred with single 3 mg Tam doses at earlier stages (E9.5-13.5; Fig. 1D,E,F,G). For adults, three 3 mg Tam doses over 6 days resulted in ~60-80% recombination of R26RlacZ or R26REYFP (supplementary material Fig. S3B; Fig. 3E-J). Ptf1aCreERTM-induced R26RlacZ recombination
in adult pancreas was Tam-dose-dependent; labeling efficiency was enhanced moving from 1 to 3 mg (supplementary material Fig. S3C).
Ptf1aCreERTM is therefore valuable for pulsed-labeling studies, and conditional inactivation of genes involved in pancreas development/cancer, with excellent spatiotemporal control.

Early Ptf1a-expressing cells contribute to all three pancreatic lineages

Ptf1aCreERTM,R26REYFP embryos were generated at various stages (Fig. 1A). Single 3 mg Tam dosing of Ptf1aCreERTM,R26REYFP pregnant mice at E10.5 would label Ptf1a-expressing cells of the primary (1°) transition (E9.5-12); pulses between E12.5-14.5 would label 2° transition Ptf1a-expressing cells; and at E15.5 or later would test whether Ptf1a-expressing cells had completely switched from multipotential behavior (Fig. 1B).

Pancreata were analyzed near the end of gestation (E18.5) for co-localization of EYFP (derivation from Ptf1a+ cells) with endocrine, ductal and acinar markers. The percentage of labeled (EYFP+) cells expressing each marker was quantified by a strategy taking into account the potential variation in proliferation and cell sizes in the acinar, duct or endocrine compartment. Because nuclear CreER localization starts ~6-12 hours post-Tam-injection with a 36-hour end point (supplementary material Fig. S2) (Ahn and Joyner, 2004; Danielian et al., 1998), data are presented as each injection point having a labeling period offset by 12-36 hours.

Tam pulses at E10.5 (E11 labeling; n=4) labeled EYFP+ cells equivalently in all three epithelium-derived compartments – acini/duct/endocrine – indicating that the Ptf1a+ population is multipotent at this stage (Fig. 1D,D′), and consistent with previous studies showing Ptf1a marks early pancreatic bud MPCs (Kawaguchi 753).
et al., 2002; Burlison et al., 2008). The percentage of EYFP+ cells per compartment was reproducibly comparable (Fig. 1D); supplementary material Fig. S1B), suggesting that the early MPC population contributes equally to all three lineages, including all endocrine cell types (supplementary material Fig. S1F).

**Ptf1a**+ MPC numbers decrease dramatically during the 2° transition

To determine the degree to which the Ptf1a+ population maintains multipotency over the 2° transition (E12.5-15.5), single Tam doses were given at E12.5, E13.5 and E14.5, with analysis at E18.5. Labeling at E13 and E14 yielded lineage-traced endocrine, duct and acinar cells (Fig. 1.E,E’,F,F’), indicating that significant numbers of Ptf1a+ MPCs exist during the 2° transition. Quantitation showed a tendency to produce acinar over duct and endocrine cells (E13, n=3; E14, n=5; Fig. 1.E,F; supplementary material Fig. S1C,D,E). In particular, the E13 result suggests that the tip epithelium at this stage already contains different populations of Ptf1a-expressing cells, in accordance with immuno-colocalization data below. These findings largely agree with CpaICreERT2 studies demonstrating diminution of the MPC pool during the 2° transition (Zhou et al., 2007). With labeling at E15, lineage-trace was predominantly restricted to acini and to a much lesser extent duct cells (n=3; Fig. 1G-G’). Taken together, these data indicate that Ptf1a-expressing cells have almost completely switched to unipotential acinar progenitor behavior by ~E15, with a minimal late-stage-remaining contribution to the duct compartment.

Shorter-term lineage-tracing was performed to assess lineage allocation of Ptf1a-expressing MPCs during the 2° transition, specifically to determine the extent to which tip MPC yield progeny moving into the trunk epithelium (the presumptive location of the endocrine/duct bipotential pool). Twenty-four hours post-labeling (injecting at E12.5, E13.5; analyzing at E13.5 or E14.5), numerous CpaI+ proacinar/acinar cells were EYFP+ (of total CpaI+ tip cells: ~8.3±2.2% at E13.5, ~11.3±2.70% at E14.5; n=4; Fig. 2.B,E,F,I). EYFP+ Hnf1β+ cells in trunk epithelium were also regularly observed but at a relatively low frequency (of total Hnf1β+ trunk cells: ~2.1±0.9% at E13.5, ~1.5±0.9% at E14.5; n=4; Fig. 2.D,E,H,I). Therefore, Ptf1a-lineage cells could contribute to both tip, and to a lesser extent, trunk compartments at ~E13/14, the stage of progression into the secondary transition proper. EYFP+Ngn3+ endocrine progenitors and EYFP+Synaptophysin+ endocrine cells (synaptophysin labels endocrine precursors and differentiated hormone+ cells), although rare in number, were found 24 hours post-labeling [of total (Ngn3+ +Synaptophysin+) cells: 0.2±0.1% at E13.5, ~0.06±0.04% at E14.5; n=4; Fig. 2.C,E,G,I]. These analyses provide direct evidence, consistent with the longer-term tracing above, that the Ptf1a-expressing tip population is already undergoing, near the initiation of the 2° transition, significant restriction of lineage multipotency. We next questioned to what extent the Ptf1a-expressing tip domain comprised heterogeneous populations of qualitatively distinct cell-types, based on their production of TFs associated with multipotency.

**2° transition tip epithelium heterogeneity: putative Ptf1a**+Sox9+Hnf1β+ MPC versus Ptf1a**+Sox9**+Hnf1β+ proacinar progenitors

We explored the signature of putative Ptf1a+ MPCs between E11.5 and E15.5 to try to move beyond the general signature of Pdx1+Ptf1a+CpaI+ cMyecreHI proposed by Zhou et al. (Zhou et al., 2007). We monitored co-localization and relative levels of Ptf1a with Sox9 and Hnf1β, two TFs having loss-of-function evidence of important roles in MPC formation (Seymour et al., 2007; Haumaire et al., 2005). At E11 and E11.75, with the initiation of tip-trunk compartmentalization, the pancreatic epithelium displayed significant heterogeneity, with three populations observable: (1) Ptf1a+BBSox9+ cells located in peripheral regions of the branching bud epithelium; (2) Ptf1a+Sox9+ cells concentrated in the prospective trunk domain; and (3) Ptf1a+Sox9+ cells scattered throughout the epithelium (Fig. 2.K).

By E12.5, tip and trunk domains were relatively distinct based on Ptf1a and Sox9 production, respectively (Fig. 2.L). Between E12.5 and E15.5, Ptf1a+Sox9+ tip cells were found distributed at both tip-trunk interface and intra-tip regions, intermingled with the Ptf1a+BBSox9+Sox9+ (proacinar) population (Fig. 2.L-O; supplementary material Fig. S4A-F). The numbers of these Ptf1a+Sox9+ cells decreased progressively over this period. Conversely, the Ptf1a+BBSox9+ population increased to comprise the entire tip domain by E15.5. Ptf1a+Sox9+ tip cells were also Hnf1β+ and CpaI+ (supplementary material Fig. S4A-F), and thus may represent the MPCs persisting during the early 2° transition, as marked by our lineage-tracing (Fig. 2P). Notably, the relative signal intensity of Ptf1a, Sox9 and Hnf1β in some of the tip MPC population seems lower compared to the robust signals in the trunk (for Sox9/Hnf1β or proacini (for Ptf1a). But, determining if lower levels of these co-expressed TFs are necessarily associated with MPC character awaits future tools that allow isolation of this population. Our co-expression and lineage-tracing data fit the notion that multipotent Ptf1a+ tip populations diminish rapidly during the 2° transition as they convert to Ptf1a+ proacinar cells.

**Ptf1a**+ cells produce only acini after birth

From birth until weaning, islet cell mass increases parallel to body mass in mouse and rat (Bouwens and Rooman, 2005). We found no evidence that endocrine cells arise from Ptf1a+expressing cells during this endocrine expansion period. Single 3 mg Tam doses were given to Ptf1aCreERT2;R26R-EYFP pregnant mice near the end of gestation (E18.5) and pancreata were analyzed at postnatal day 21 (P21; n=4; Fig. 3A). EYFP was found exclusively in CpaI+ acinar cells (Fig. 3B-D), suggesting that postnatal Ptf1a-expressing cells contribute to neither endocrine nor duct compartments. To determine if duct or endocrine cells arise from Ptf1a+ cells during normal homeostasis in young adults, we treated P30 mice with three doses of Tam (Fig. 3A). Two days after the last injection (P36), a high proportion of acinar cells were labeled (60.9±3.83%; n=4; Fig. 3E-G), increasing to 72.6±5.39% after 6 months post-Tam. No EYFP+ cells were found in the duct or endocrine compartments at either age (P36, 7-month; n=4; Fig. 3H-J). We also failed to detect Hnf1β+Ck19+ (Ck19 is also known as Krt19 – Mouse Genome Informatics) centroacinar cells (CACs) labeled with EYFP after 6-month chase (Fig. 3K,K’L,M), consistent with a lack of Ptf1a immunoreactivity in Sox9+Ck19+ and Hnf1β+Ck19+ CACs (Fig. 3M’-M”; supplementary material Fig. S6A). These latter results suggest that Ptf1a expression does not mark CACs in vivo, contrasting a previous study reporting Ptf1a expression in flow-sorted CACs (Rovira et al., 2010). Together, these data suggest that Ptf1a+ cells self-replicate to maintain the acinar pool in the adult organ, with no contribution towards the duct or endocrine populations.

**PDL induces ductal transdifferentiation of Ptf1a**+ acinar cells, reactivating MPCs and endocrine progenitor factors

We endeavored to determine whether Ptf1a-expressing acinar cells could exhibit facultative progenitor activity under PDL, exhibiting
Fig. 2. Ptf1a⁺ tip cell contribution to trunk and endocrine progenitor/precursor compartment, and heterogeneity in 2°-transition tip epithelium (with putative Ptf1aLO MPC and Ptf1aHI proacinar cells). (A) Short-term lineage-tracing schematic. (B-D, F-H) EYFP-labeled CpaI⁺ tip cells, Hnf1β⁺ trunk cells, and Ngn3⁺Synaptophysin⁺ (Syn) endocrine precursors were detected 24 hours post-labeling of Ptf1a-expressing cells at (B-D) E12.5 or (F-H) E13.5. (E-I) Percentage EYFP⁺ cells in each lineage with labeling at E12.5 (E; n=4) or E13.5 (I; n=4). (J-O) Ptf1a⁺Sox9 immunodetection between E11-15.5 showing tip epithelium with two Ptf1a⁺ populations: Ptf1a⁺Sox9⁺ putative MPC (arrowhead), and Ptf1a⁺Hnf1β⁺ proacinar progenitors. Numbers of Ptf1a⁺Sox9⁺ cells decreased rapidly as organogenesis progressed. (P) Schematic, lineage potency of Ptf1a⁺Sox9⁺Hnf1β⁺ tip MPC during the 2° transition. Scale bars: 50 μm.
multipotentiality similar to embryonic Ptf1a-expressing MPCs. PDL was reported to induce fairly efficient de novo generation of facultative Ngn3+ endocrine progenitors (Xu et al., 2008), and this model was chosen to evaluate whether Ptf1a+ acinar cells could convert toward the Ngn3+ population. Five-week-old Ptf1aCreERTM;R26REYFP mice given three doses of Tam had PDL performed 2 weeks after the last Tam injection, as described (Xu et al., 2008) (Fig. 4A). An average ~60-80% labeling efficiency was observed and strictly restricted to acinar cells (Fig. 3J); high labeling efficiency was deliberately pushed so as to detect minor populations of facultative progenitors. Pancreata were analyzed 5 days (PD L D5), 1 week (PD L D7), 4 weeks (PD L D30) or 8 weeks (PD L D60) post-PDL.

The ligated distal part of the pancreas (hereafter ‘PD L tail’), which consists of ~80% of the splenic lobe, underwent massive acinar regression by PD L D7 (supplementary material Fig. S5B,D). The unligated proximal pancreas (hereafter ‘PD L head’, comprising ~20% of the remaining unligated splenic lobe, intact gastric lobe and ventral pancreas) remained unchanged morphologically after ligation and served as a control for labeling. 

| Fig. 3. Ptf1a-expressing cells are acinar-restricted after birth. (A) Postnatal/adult lineage-tracing schematic. (B,C) Ptf1aCreERTM;R26REYFP embryos Tam-pulsed at E18.5, analyzed 3 weeks after birth showing EYFP exclusively in acini. (D) P21; percentage of each pancreatic cell type expressing EYFP. (E,F,H,I) 1-month-old Ptf1aCreERTM;R26REYFP mice treated with Tam were chased for (E,F) 1 week or (H,I) 6 months; EYFP was only in Cpa1+ acinar cells. (G,J) Comparing EYFP+ traced cells at P36 (G) or 7 months (J) showed similar acinar labeling, suggesting homeostatic acinar self-replication. (K-L) EYFP was absent from (K,K') Ck19+ or (L) Hnf1β+ CAC cells after 6-month chase; diagrammed as Ptf1a-expressing cells not producing CACs. (M-M′′′) Ptf1a protein was undetectable in Sox9+Ck19+ CACs. Scale bars: 50 μm. |
Fig. 4. PDL-induced acinar-to-ductal transdifferentiation and de novo Ngn3 activation. (A) PDL lineage-tracing schematic. (B-F) At D5, Ptf1a protein was found only in acini in (B) sham tail and (C) PDL head, but was in (D) PDL tail duct cells. Ngn3 was found only in islets of (E) sham tail or (F) PDL head. (G) De novo activation of Ngn3 in Ck19+ duct cells, PDL tail. (H) Quantification, average number Ngn3+Ck19+ duct cells per section, PDL D7 and D30 (~10-15 sections entirely counted per PDL tail; n=3); solid line indicates mean. (I,J,M,N) EYFP+Ngn3+ cells were absent from sham tail and PDL head at (I,J) PDL D7 or (M,N) D30. (K,O) A small fraction of the Ngn3+Ck19+ duct cells were derived from Ptf1a-lineage-labeled cells (EYFP+, arrowhead) in PDL tail at (K) PDL D7 and (O) D30. (L,P) Quantification of (L) percentage of EYFP+Ck19+ cells that were also Ngn3+, and (P) percentage of EYFP+Ngn3+Ck19+ cells over total Ngn3+Ck19+ cells at PDL D7 and D30. (Q,R) EYFP+Ck19+Hnf1β+ duct cells were absent from (Q) sham tail or (R) PDL head. (S) Ptf1a-lineage-derived ducts (EYFP*Ck19+) were Hnf1β+ (arrowhead) in PDL tail at PDL D7, indicating acinar-to-ductal transdifferentiation. Scale bars: 50 μm.
efficiency (supplementary material Fig. S5C). After PDL, the total pancreatic epithelial area of PDL tail was dramatically reduced, while overall pancreatic weight remained similar because acini were replaced by fibrotic and inflammatory cells; islets and duct cells were spared and the animals remained euglycemic (supplementary material Fig. S5C-H) (Xu et al., 2008). Post-PDL D5, tubular complexes composed of Ck19+ ducts started to form in the PDL tail. Ptf1a protein was detected in these tubular complexes at this stage but not in the ducts of the PDL head, or the control sham-operated tail (hereafter ‘sham tail’; Fig. 4B-D; supplementary material Fig. S5A). The presence of Ptf1a protein in ductal cells proximate to the regressing acini might reflect detection of the early stages of acinar-to-ductal conversion. An alternate explanation is that remodeling ducts activated Ptf1a expression; a hypothesis ruled out by two observations: (1) the lack of Ptf1a protein by immunolabeling in the Ck19+/Hnf1β+ tubular complexes at PDL D7, D30 and D60 (supplementary material Fig. S6B-M); and (2) Tam treatment of Ptf1aCreERT2;R26R-EYFP/EYFP mice post-PDL D7, after acini involution was almost completed, such that CreER would only label duct cells activating Ptf1a expression. By this strategy, extremely rare Ptf1a-lineage-labeled duct cells (~1-2 EYFP*Ck19+ every three sections, possibly derived from late incorporation into the tubular complexes of the 1% acinar cells that escape involution) were found at post-PDL D30 and D60 (supplementary material Fig. S7E-K). Taken together, these results indicate that duct cells did not activate Ptf1a expression.

One week post-PDL, ~99% of acinar cells in the PDL tail had invovluted, and the remodeled ducts formed highly proliferative tubular complexes. The PDL tail was fibrotic and infiltrated with inflammatory cells (supplementary material Fig. S5A-D). As previously reported (Xu et al., 2008), we detected Ngn3 protein in the islet endocrine cells (supplementary material Fig. S5A-D). To assess whether Ptf1a+ cells can give rise to endocrine cells after injury, particularly focusing on β-cells, pancreata from Ptf1aCreERT2;R26R-EYFP/EYFP mice lineage-labeled by Tam treatment prior to PDL were analyzed, first at post-PDL D30 (Fig. 5A). In order to increase the sensitivity of detecting a potentially small population of endocrine cells derived from Ptf1a+ facultative progenitors, we performed an extensive, systematic analysis across a large portion of each pancreas. Thus, we analyzed ~40-45 sections from each mouse (~10-15 sections per ~10-15 sections counted per PDL tail pancreas) (n=3; Fig. 4H). Scattered rare Ngn3+Ck19+ duct cells were also found at post-PDL D60 (not shown), consistent with the previous post-PDL analysis showing Ngn3 mRNA expression peaking at post-PDL D7 and reducing by later time points (Xu et al., 2008). These data also imply that the majority of the early Ngn3+Ck19+ protodendocrine cells have moved on to another differentiation state at later time points post-PDL. Notably, the Ngn3 signal was much higher in duct cells than in the islet endocrine cells (Fig. 4F,G, inset). These data imply that while Ngn3 is required for proper function/maintenance of adult endocrine cells (Wang et al., 2009), Ngn3+ duct cells arose from de novo activation of endocrine progenitor behavior upon injury, resembling the embryonic endocrine neogenesis process.

To determine whether acinar cells contribute to regeneration of the duct and endocrine pancreas after PDL, we used Ptf1aCreERT2 to lineage-trace the Ptf1a+ acinar cells and their progeny. In PDL tail post-PDL D7, we observed small numbers of EYFP+ cells in Ck19+ ducts, suggesting that some of the Ptf1a-lineage-labeled acinar cells escaped involution but became integrated into the tubular complex (Fig. 4Q-S; supplementary material Fig. S7A-D; Fig. S8F-H). The number of EYFP*Ck19+ duct cells increased substantially from post-PDL D7 to D60 (from 5.8±1.2% to 25.4±5.14% of total Ck19+ duct, respectively; n=3; supplementary material Fig. S7A-D, A’-C’), consistent with a continued high proliferative capacity as detected by BrdU incorporation (supplementary material Fig. S5E-G, I).
throughout the tissue and accounted for 25.4±5.14% of the total Ck19+ duct area. This representation was 11.6±1.81% at post-PDL D30, suggesting that proliferation and expansion of the Ptf1a-lineage-labeled duct/tubular-complex cells continued from D30 to D60 (n=3; supplementary material Fig. S7B-D). We also detected in post-PDL D60 tissue a small but significant number of EYFP+endocrine-hormone+ cells in Ck19+ ducts (Fig. 5H), as well as in islets (Fig. 5I). Around 75% of the EYFP+hormone+ cells were located in/at the ducts, whereas ~25% were found in the islets. Some of these endocrine cells, either in the ducts or in islets, were also insulin-producing (Fig. 5J,M), suggesting that β-cells can be derived from Ptf1a+ acinar cells after PDL. Most importantly, these Ptf1a-lineage-derived insulin+ cells produced several mature β-cell transcription factors, such as Pdx1HI (Fig. 7A,B), Nkx6.1 (Fig. 7E,F), and MafA (Fig. 7J,L), indicating a mature β-cell status. Most of the EYFP+insulin+ cells were presents as singlet β-cells located at/in the ducts and did not produce GLUT2 (data not shown). The lack of β-β cell contact may prevent membrane localization of GLUT2 in these EYFP+ singlet β-cells as previously suggested (Orci et al., 1989; Gannon et al., 2000). Extensive, systematic quantification revealed 965±393 EYFP+hormone+ endocrine cells, per PDL tail analyzed (n=4; Fig. 5N). We only observed one to eight islets per 10% of PDL tail containing EYFP+ endocrine cells; in such islets only ~5-10% of the cells were EYFP+ (n=4; Fig. 5O; data not shown). Because ~99% of acinar cells were lost at post-PDL D7, and the persistent...
blockage of the pancreatic duct limits acinar regeneration, we believe that these EYFP+ endocrine cells were originated from Ptf1a+ acinar-derived duct cells, rather than arising directly from Ptf1a+ acini as no amylase+ insulin+ transitional intermediate was found at PDL D7, D30 and D60 (supplementary material Fig. S9C-H). We did not detect any EYFP+ cells in the endocrine compartment for either sham tail or PDL head at post-PDL D60 (Fig. 5F,G,J,K,N,O). These in vivo lineage-tracing data support the proposal that Ptf1a+ acinar cells could be stimulated to become facultative progenitors and undergo a long-term (albeit inefficient) reprogramming to endocrine cells, via a ductal intermediate, in the PDL-injury/regeneration context (Fig. 5P).

Enhanced acinar-to-duct-to-endocrine conversion with pre-existing β-cell ablation

Because islet cells were spared and mice remained euglycemic after PDL, we propose that the lack of hypothetical physiological stimuli to produce additional β-cells could explain the low level conversion to endocrine cells from Ptf1a+ acini-derived duct cells. We reasoned that eliminating some pre-existing β-cells could provide one such physiological cue (perhaps including the ensuing hyperglycemia), and addressed the deep injury context of PDL plus β-cell destruction. Tam-treated Ptf1aCreERTM;R26REYFP/EYFP mice were given two consecutive doses of STZ three days post-PDL to kill a portion of β-cells, and pancreata were analyzed 30 and 60 days post-PDL (Fig. 6A). We observed ~50-70% reduction in β-cell numbers with this protocol, and mice became hyperglycemic (ad lib feeding blood glucose level >450 mg/dl) 2-3 days after STZ treatment (supplementary material Fig. S10B).

To determine if [PDL+STZ] treatment could speed up acinar-to-endocrine conversion, we analyzed pancreata post-PDL D30. [PDL+STZ]-treated mice remained hyperglycemic at this time (supplementary material Fig. S10B). We did not treat mice with any β-cell-proliferation-stimulating agents (e.g. insulin, GLP or...
EGF/gastrin), under the logic that avoiding such agents might promote transdifferentiation and disfavor replication of \( \beta \)-cells. We found EYFP+endocrine-hormone+ cells mostly located either in ducts, or duct-apposed, in [PDL+STZ] tail tissue already by post-PDL D30, notably different from PDL alone at D30 (Fig. 6D; for more examples see supplementary material Fig. S11A). No EYFP+endocrine-hormone+ cells were found in the PDL head or STZ-treated sham tail (Fig. 6B,C), suggesting strongly that STZ alone did not induce acinar-to-endocrine transdifferentiation. The numbers of EYFP+endocrine-hormone+ cells found in [PDL+STZ] D30 tail tissue were similar to those found post-PDL D60 tail without STZ treatment (average 990 ± 149 EYFP+endocrine-hormone+ cells per PDL tail in [PDL+STZ] D30, compared to 965 ± 393 EYFP+endocrine-hormone+ cells per PDL tail at PDL D60; n = 3 separate animals; compare Fig. 6N with Fig. 5N). At post-[PDL+STZ] D60, the numbers of EYFP+endocrine-hormone+ cells had increased ~3-4.5 fold over D30 (average 3653 ± 1205 EYFP+endocrine-hormone+ cells per PDL tail tissue; n = 3; Fig. 6N). EYFP+insulin+ cells were also found at post-[PDL+STZ] D30 and D60 (Fig. 6L,M; see more examples in supplementary material Fig. S11A,B). These EYFP+insulin+ cells were also Pdx1HI (Fig. 7C,D), Nkx6.1+ (Fig. 7G,H), and Mafa+ (Fig. 7K,L), suggesting that they are mature \( \beta \)-cells. Notably, fewer EYFP+endocrine-hormone+ cells were located within islets under the [PDL+STZ] condition compared to PDL alone at day 60 (average ~1 islet per 10% PDL tail labeled in post-[PDL+STZ] D60, versus average ~5 islets per 10% PDL tail in post-PDL D60; compare Fig. 5O; Fig. 6O). We hypothesize that repellents or toxics released from apoptotic \( \beta \)-cells after STZ treatment might render the islets unattractive to home Ptf1a-lineage-derived endocrine cells. Taken together, these findings demonstrated that the deep lesion enhanced acinar-to-endocrine transdifferentiation rate and efficiency compared to PDL alone.

**DISCUSSION**

We are still far from understanding how dynamic cell heterogeneity and intercommunication connect to epithelial morphogenesis and cell-fate allocation, and if certain mature cells retain homeostatic or facultative multipotency/plasticity. Our immuno-colocalization of TFs considered as progenitor-defining revealed heterogeneity as epithelial morphogenesis begins at E11.5-12.5, well before the 2° transition stage. The putative Ptf1a+Sox9+Hnf1\( \beta \)+ MPCs (the same signature as early MPCs), possibly under the influence of localized signals from the mesenchyme, become localized to the tip epithelium and progressively diminish between E12.5 and E15.5. The reduction in these putative MPCs parallels the quantitative decrease in progeny of Ptf1a-expressing cells that enter non-acinar lineages. After the 2° transition (E15.5 onward), Ptf1a+ MPC activity becomes very low, and Ptf1a-expressing cells are acinar-unipotent by late gestation and in adults.
More importantly, we showed that PDL or [PDL+STZ] induced restoration of an early-organogenesis-like MPC competence in adult acinar cells, with probably stepwise redifferentiation into ducts then endocrine cells. Ptf1α+ acinar cells reactivated embryonic multipotency markers, differentiating rapidly into long-lived duct cells. Some Ptf1α-derived duct cells activated Ngn3, and some Ptf1α-lineage-derived cells progressed to endocrine cells, including Pdx1+/Nkx6.1+/MafA/insulin−β-cells. This conversion occurred without exogenous factors such as β-cell-instructive TFs or intercellular signaling factors.

**Progressive lineage restriction of Ptf1α-expressing cells**

We have more carefully enumerated the multi-to-unipotential switch during organogenesis, with findings largely consistent but with novel aspects compared to the previous CpaKCre study (Zhou et al., 2007), but it is also important that we used an MPC-instructive gene Ptf1α, and not a Ptf1α surrogate encoding a digestive enzyme, which should have no regulatory role in MPC formation/maintenance. Furthermore, our E10.5 immunocolocalization data (supplementary material Fig. S12) showed Cpa in ~65% of Ptf1α− cells (all Cpa− cells were Ptf1α−), suggesting Cpa only partially marks MPC. Our Ptf1αCreERTM-based analysis validates the CpaKCre study but elaborates a stronger understanding of the dynamic nature of MPC location during tip-trunk segregation in the 2° transition.

The CpaKCre study proposed that Cpa+ tip MPC contribute significantly to the expansion of the trunk epithelium (endocrine/duct bipotent progenitor compartment) after E12, but we found that Ptf1α-expressing cells seeded the trunk epithelium until E13.5, with only a small contribution at/after ~E12.5. We can conclude that tip seeds trunk is a minor influence during the true 2° transition, consistent with the proposal that intra-trunk proliferation and plexus remodeling (Villasenor et al., 2010) cause most of the growth and extension of the trunk epithelium.

We note that the Ptf1α′Pdx1′Cpa+eMycHI MPC proposed in Zhou et al. (Zhou et al., 2007) did not refer to tip heterogeneity, and in fact differentiated acini retain that signature. Solar et al. (Solar et al., 2009) suggested that Hnf1β− trunk cells are multipotent between E11.5-13.5, but did not rule out that Hnf1βCreERTM-traced MPC could be an Hnf1β− Cpa+ tip subset. Furthermore, considering the tip-trunk interface Nkx6.1+Ptf1α′Sox9′Pdx1′ MPC proposed by Kopp et al. (Kopp et al., 2011), which lacked Hnf1β, the Ptf1α′Sox9′putative tip MPC we detected were Hnf1β+, and distributed heterogeneously, both intra-tip and at the tip-trunk interface. The Ptf1α′Sox9′Hnf1β+ tip MPC could be the same as those detected by Solar et al. (Solar et al., 2009) and Kopp et al. (Kopp et al., 2011), leading to the generalization that multipotentiality is preserved in the tip, not trunk, domain. What controls tip-trunk compartmentalization, and which signals are present in the local tip microenvironment that determine this MPC niche, represent interesting aspects for future studies.

**Postnatal lineage restriction of Ptf1α+ cells**

Postnatal Ptf1α+ cells only produced acinar cells, agreeing with the results of ElastaseCreERT2 studies showing homeostatic maintenance of this compartment comes from acinar replication (Desai et al., 2007). A Sox6CreERTK knock-in study (Furuyama et al., 2011) seems contradictory, in suggesting that ~80% of acinar cells came from Sox9+ duct or centroacinar cells following a one-year post-Tam chase. We found no significant reduction in the proportion of Ptf1αCreERTM-labeled acini after a 6-month chase, arguing against dilution from other cell populations. This discrepancy could be reconciled, bearing in mind the lower level Sox9 expression in adult acini (Kopp et al., 2011; Dubois et al., 2011) (Fig. 3M,M∗), such that the high tamoxifen dosage used (5×4 mg) in Furuyama et al. (Furuyama et al., 2011), would broadly and directly activating the reporter allele in acini.

**Facultative PDL-induced MPC behavior in adult acinar cells**

The mass of acinar cells and their relatively close developmental history with endocrine cells lead to ideas that partial reprogramming in vivo might offset a β-cell deficit without compromising organ function (Pan and Wright, 2011). Zhou et al. (Zhou et al., 2008) reported reprogramming of acini to β-like fate using a viral cocktail of β-cell TFs: Pdx1, Ngn3 and MafA. Our evidence suggests that Ptf1α+ acinar cells possess facultative progenitor potential under PDL, allowing them quickly to produce long-lived ducts and later-ariising endocrine/β-cells. Several *in vitro* studies suggested acinar-to-β-cell conversion with EGF, nicotinamide, TGFα or combination(s) treatment (Baeyens et al., 2005; Minami et al., 2005). Non-autonomous effects of tissue dissociation might substantially affect *in vitro* conversion. Genetic lineage-tracing was not included in the previous reports of *in vivo* acinar-to-endocrine transdifferentiation (purportedly involving amylase−insulin intermediates) in 24-hour post-PDL rats (Bertelli and Bendayan, 1997). Our study therefore provides rigorous *in vivo* evidence for PDL-induced acinar-to-endocrine conversion. We contradict the conclusion from ElastaseCreERT2 lineage-tracing *in vivo* that acini did not seed duct or endocrine fates after Ppx, PDL and cerulean-induced pancreatitis (Desai et al., 2007), but the 30% acinar labeling efficiency (compared to our ~80% labeling, assessing large amounts of the organ and over a long period) might have failed to detect low-level contributions.

If there is any cryptic destabilization of the acinar program from Ptf1α heterozygosity (Ptf1αCreERTM is a null allele), there are consequences only under injury, and Ptf1αCreERTM− mice have normal embryonic and adult pancreas. Distinct Ptf1α levels might control the interaction with the RBP-J or RBP-JL partner in the PTF1 complex. Ptf1α associates with RBPJ (PTF1RBP) in MPC, but with RBPJL (PTF1RBPβ) in proacinar/acinar cells (Masui et al., 2007), with the PTF1RBPα to PTF1RBPβ switch likely essential for facilitating epithelial tip-trunk compartmentalization (Masui et al., 2007; Schaffer et al., 2010). Under injury, reduced Ptf1α levels could allow reformation of PTF1RBPβ, which together with Sox9/Hnf1β production leads to reacquisition of MPC character. Future studies will be directed toward understanding how differential Ptf1α levels affect multipotency restoration in Ptf1α+ acini during injury.

Acinar-to-endocrine conversion could be aided by disturbances in the extracellular environment, including the chronic inflammation-associated signals in PDL. Cytokines may also influence pancreas regeneration (Homo-Delarche and Drexhage, 2004), and invading macrophages may suppress islet loss during exocrine degeneration (Tessem et al., 2008). Wang et al. (Wang et al., 2007) suggested a role for immune-reaction effects against adenovirus in liver-β-like cell conversions driven by pro-endocrine TFs (Pdx1, Ngn3). Our Ptf1α-lineage-labeled endocrine cells were often found next to duct-apposed small endocrine clusters. Such clusters are distributed throughout PDL pancreas, although some might arise from duct cells proper (Xu et al., 2008). Speculatively, acinar-derived endocrine cells could home to these small endocrine clusters rather than to islets proper, or endocrine...
cell migration from their birthplace in the tubular complexes might be hindered by the altered fibrotic stroma.

While we showed that Ptf1a+ acinar cells produced Ck19\Hnf1β\Sox9+ primitive ductal intermediates, Ngn3+ endocrine progenitors and hormone+ endocrine cells, showing directly a sequential movement through these transitional states would need either real-time tracing, or dual/multiplex genetic lineage-tracing methods (e.g. combining Cre/LoxP, Dre/Rox or FLP/FRT). We did not observe any pan-endocrine/insulin co-label with acinar markers arguing against a direct rapid conversion from acini to endocrine cells.

Reproliferative CACs have been suggested as facultative progenitors under injury. Rovira et al. (Rovira et al., 2010) detected Ptf1a by RT-PCR in Aldh1+ CACs, and MPC competence in CAC-derived pancreatospheres. We failed to detect Ptf1a protein in CACs post-PDL, which incorporated into tubular complexes and yielded duct/endocrine cells. However, Tam-induction at post-PDL in CACs post-PDL, which incorporated into tubular complexes and derived pancreatospheres. We failed to detect Ptf1a protein in endocrine progenitors and hormone + endocrine cells, showing Acinar to endocrine reprogramming captured via their induced growth potential; indeed, endocrine conversion might increase under strategies similar to those applied to iPSCs. Our findings might lead to pharmacological intervention using small molecules that modulate the epigenetic landscape or mimic inflammatory signaling. Controlling plasticity might allow regulated reprogramming as an alternative to cell transplantation.

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Competing interests statement
The authors declare no competing financial interests.

Supplementary material
Supplementary material available online at http://dev.biologists.org/suppl/doi:10.1242/dev.090159/-DC1

References


Fig. S1. Lineage tracing of Ptf1a-expressing cells using the novel Ptf1aCreERTM allele. (A) Schematic representation of Ptf1aCreERTM knock-in generation via recombinase-mediated cassette exchange (RMCE). A Ptf1aCreERTMHygroR cassette exchange vector was constructed by inserting a 4105 bp fragment of Ptf1a into a plasmid containing two inversely oriented LoxP sites, and a 5' UTR NotI site was changed to FseI. DNA encoding CreERTM (a gift from Andrew McMahon, Harvard) was then inserted between this FseI and a natural MluI site in Ptf1a exon 1. A pgk-driven hygromycin resistance gene (HygroR), flanked by tandem FRT sites, was inserted at the 3' end of the exchange vector for positive selection during RMCE. RMCE was as previously described (Long et al., 2004). An embryonic stem (ES) cell clone 5D12 containing the Ptf1aLCA (Burlison et al., 2008) was electroporated with the Ptf1aCreERTMHygroR exchange vector and a Cre vector. Clones surviving hygromycin/gancyclovir selection were screened using PCR (primers sequences available upon request). Chimeric mice derived from injecting clone 5D12:1D7 ES cells into C57BL/6J blastocysts were bred with C57BL/6J mice to derive Ptf1aCreERTMHygroR. (B-E) The HygroR cassette was removed by FLPe deleter mice (provided by Susan Dymecki, Harvard). Scatter plots showing the percentage of cells in each pancreatic compartment expressing Eyfp after Tam injection at E10.5 (B; n=4), E12.5 (C; n=3), E13.5 (D; n=5), and E14.5 (E; n=3). Each point represents a single pancreas. (F) Ptf1a-expressing cells at E11 contribute equally to all endocrine cell types; n=3. A, acinar; D, duct; E, endocrine.
Fig. S2. Cre production recapitulate endogenous Ptf1a production. (A) Cre protein is located in the cytoplasm of Ptf1a⁺ cells in the absence of tamoxifen at E16.5 (upper panel); tamoxifen treatment resulted in nuclear translocation of Cre in Ptf1a⁺ cells (lower panel). (B) Cre protein was not detected in wild-type E16.5 CpaI⁺ cells, and tamoxifen treatment caused nuclear localization of Cre in CpaI⁺ cells. Scale bar: 50 μm.
Fig. S3. Ptf1a<sup>CreERTM</sup>-induced recombination at the ROSA26<sup>R<sub>lacZ</sub></sup> allele is Tam and Tam-dosage dependent. (A) Schematic presentation of Ptf1a<sup>CreER</sup>-mediated recombination at the Rosa26<sup>Eryfp</sup> locus only in the presence of Tam. (B) Whole-mount ∝-gal-stained adult gut tissues showed that ∝-gal staining was found in the pancreas only in the presence of Tam (left panel), but not in corn-oil injected control (right panel), indicating the absence of leakiness. (C) There was a Tam-dosage-dependent increase (from 3 to 9 mg) in the ∝-gal<sup>+</sup> cell numbers in the pancreas. (D) Efficient embryonic labeling at E16.5 after a single dose 3 mg Tam injection (left panel). ∝-gal staining was mainly found in acinar cells. ∝-gal staining was not found in the pancreas of mice that did not carry the Ptf1a<sup>CreERTM</sup> allele (B, middle panel; D, middle panel).
Fig. S4. Distribution of Ptf1a^+Sox9^+Hnf1β^+ tip MPCs during the 2° transition. Immunolabeling analysis of Ptf1a, Sox9 and Hnf1β at (A) E12.5, (B) E13.5, (C) E14.5 and (D) E15.5 shows the distribution and decreasing numbers of Ptf1a^+Sox9^+Hnf1β^+ tip MPCs (arrowhead and arrow) in both intra-tip and tip-trunk interface region. A definite intra-tip location is defined by the location at or near the most distal tip region of the pancreatic epithelium, contact with adjacent pancreatic mesenchyme, and importantly the absence of any cells positive for trunk markers (Sox9^+Hnf1β^+) next to that cell’s location, from analysis of all of the adjacent sections (arrowhead). ‘Tip MPCs’ that are located at the tip-trunk interface are defined by proximity to the Sox9^+Hnf1β^+ trunk cells (arrow). The Ptf1a^+Sox9^+ tip MPCs are also CpaI^+ (arrowhead) at (E) E13.5 and (F) E14.5. Scale bar: 50 μm.
Fig. S5. PDL induces dramatic increased in proliferation in the remodeling ducts. (A,B) Gross morphological appearance of PDL tail at post-PDL D7 (B) compared to sham tail (A). (C,D) Sections of sham tail (C) and PDL tail (D) stained with Hematoxylin and Eosin (H&E). (E-I) Significant increased BrdU incorporation in the ducts of PDL D7 and D30 tail (F,G,I) compared with PDL head (E,I) indicate active proliferation of remodeling ducts. No significant changes in BrdU labeling in islet cells between PDL head and tail tissues (H). *P<0.005. d, ducts; I, islet; NS, not significant.
Fig. S6. Ptf1a production in mature adult pancreas and PDL-injured pancreas. (A) Ptf1a protein is not detected in Hnf1β+CK19+ CACs. Ptf1a protein is only detected in acinar cells, but not in Hnf1β+CK19+ duct cells in both PDL head and tail tissues at PDL (B-G) D7, (H-J) D30 and (K-M) D60. i, islet.
Fig. S7. Ptfla lineage-labeled duct cells were derived from acinar-to-ductal transdifferentiation, and not direct activation of Ptfla expression in the ducts post-PDL. (A-C’) Ptfla-lineage labeled (Eyfp’Ck19’) ducts in PDL tail at PDL (A,A’) D7, (B,B’) D30, and (C,C’) D60. (D) Quantitation analysis of percentage of Ptfla-derived duct cells over total Ck19+ duct cells. (E) Schematic of post-PDL tamoxifen-treatment lineage-tracing analysis. Labeling efficiency in acinar cells remained high in PDL head tissue when tamoxifen was administrated post PDL at PDL (F) D30 and (I) D60. Very rare Ptfla-derived Ck19+ duct cells were found at PDL (G,H) D30 and (J,K) D60 when tamoxifen was injected at post-PDL D7.
Fig. S8. PDL induces acinar-to-ductal transdifferentiation and activates Ngn3 protein production in duct cells. (A-B) Additional examples of Ptf1a-lineage-labeled Ngn3+Ck19+ duct cells at post-PDL D7 (A-A') and D30 (B-B'). (C-N) Ptf1a+ acini gave rise to long-lived Hnf1β+ (E) and Sox9+ (H,K) duct cells in PDL tail at post-PDL D7 and D30, but not in sham tail (C,F,I) or PDL head (D,G,J). The Sox9+Hnf1β+ duct cells are Eyfp– in sham tail (L) or PDL head (M). Most of the Ptf1a+ acinar-derived duct cells at PDL D7 are Sox9+Hnf1β+ (N). Scale bar: 50 μm.
Fig. S9. PDL did not induce direct acinar-to-endocrine transdifferentiation. (A-B') Hnf1β⁺ duct cells did not express amylase at PDL D7 in both head (A) and tail (B',B'), suggesting that acinar enzymes were downregulated before the activation of multipotency factors, such as Hnf1β. (C-H') Amylase was only found in acini in PDL head at PDL D7 (C), D30 (E) and D60 (G). No Amylase-hormone⁺ transitional cell state was found in PDL tail post-PDL D7 (D,D'), D30 (F,F') and D60 (H,H'), suggesting that Ptf1a-lineage-derived endocrine cells are not a result of rapid, direct acinar transdifferentiation, but stepwise acinar-to-duct-to-endocrine reprogramming.
Fig. S10. Strategy for tissue collection and quantitation analysis. Sham tail and PDL tail were embedded in OCT for cryosectioning. (A) The whole tissue block was sectioned, we examined ~40-45 sections, taken at 100 µm distance, and stained with a cocktail of endocrine hormones (including insulin, glucagon, somatostatin and pancreatic polypeptide). The pictures were taken at 10´ magnification to include all the hormone+ areas. The islet areas were measured using NIH ImageJ (v. 1.43.76) software. (B) PDL+STZ treated mice remained hyperglycemic at PDL+STZ D30 and D60. While there is a significant increase in Ptf1a-lineage-derived insulin+ β-cells, the small number of β-cells generated in this transdifferentiation process is therefore insufficient to improve overall glucose homeostasis.
**Fig. S11.** *Ptf1a* lineage-labeled endocrine/β-cells. Additional examples of *Ptf1a*-lineage-labeled endocrine-hormone⁺ cells (upper panel) and insulin⁺ cells (lower panel) at [PDL+STZ] D30 (A) and [PDL+STZ] D60 (B).
Fig. S12. Numerous Ptf1a+ cells do not co-express Cpal at E10.5. (A-F) Immunolabeling analysis of Ptf1a and Cpal from different regions of E10.5 dorsal pancreas bud shows that the Ptf1a+Cpal− cells (arrowhead) were frequently found at this stage, suggesting Cpal may only marks a fraction of MPC populations. Scale bar: 50 μm.
Table S1. Antibodies

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<th>Secondary antibodies</th>
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