

#### STEM CELLS AND REGENERATION

## **RESEARCH ARTICLE**

# A single-cell atlas of *de novo* $\beta$ -cell regeneration reveals the contribution of hybrid $\beta/\delta$ -cells to diabetes recovery in zebrafish

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#### **ABSTRACT**

Regeneration-competent species possess the ability to reverse the progression of severe diseases by restoring the function of the damaged tissue. However, the cellular dynamics underlying this capability remain unexplored. Here, we have used single-cell transcriptomics to map de novo β-cell regeneration during induction and recovery from diabetes in zebrafish. We show that the zebrafish has evolved two distinct types of somatostatin-producing  $\delta$ -cells, which we term  $\delta$ 1- and  $\delta$ 2-cells. Moreover, we characterize a small population of glucose-responsive islet cells, which share the hormones and fate-determinants of both  $\beta$ - and  $\delta$ 1-cells. The transcriptomic analysis of  $\beta$ -cell regeneration reveals that  $\beta/\delta$  hybrid cells provide a prominent source of insulin expression during diabetes recovery. Using in vivo calcium imaging and cell tracking, we further show that the hybrid cells form de novo and acquire glucose-responsiveness in the course of regeneration. The overexpression of dkk3, a gene enriched in hybrid cells, increases their formation in the absence of β-cell injury. Finally, interspecies comparison shows that plastic  $\delta$ 1-cells are partially related to PP cells in the human pancreas. Our work provides an atlas of  $\beta$ -cell regeneration and indicates that the rapid formation of glucoseresponsive hybrid cells contributes to the resolution of diabetes in zebrafish

KEY WORDS: Beta-cell, Insulin, Pancreas, Regeneration, Single cell, Zebrafish, Gamma-cell, Cell fate, Cell plasticity, Diabetes

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#### **INTRODUCTION**

The pancreas, a glandular organ, is composed of exocrine and endocrine compartments. The exocrine part of the pancreas is involved in food digestion via the function of acinar and ductal cells. The endocrine part is responsible for metabolic homeostasis, specifically regulation of blood glucose levels. The endocrine compartment is organized into the islets of Langerhans, which consist of three major cell types:  $\alpha$ -,  $\beta$ - and  $\delta$ -cells. The three cell types produce and secrete specific hormones: α-cells generate glucagon (gcg),  $\beta$ -cells generate insulin (ins) and  $\delta$ -cells generate somatostatin (sst). Blood glucose homeostasis is achieved by regulated secretion of endocrine hormones in the blood stream; insulin activates glucose uptake in peripheral tissue, while glucagon opposes insulin action by stimulating the production of sugar in the liver (Noguchi and Huising, 2019). Somatostatin release has an inhibitory role for both insulin and glucagon secretion, as its loss triggers acute hypoglycemia (Li et al., 2018). As the selective autoimmune destruction of β-cells causes type 1 diabetes, enhancing the limited regenerative potential of the pancreas has become an important quest towards developing new regenerative therapies.

Regeneration and cell plasticity are intimately connected. Upon near-complete  $\beta$ -cell destruction, a small proportion of glucagon-expressing  $\alpha$ -cells trans-differentiate into  $\beta$ -cells in adult mice and embryonic zebrafish (Chera et al., 2014; Thorel et al., 2010). In addition, a subset of terminal duct cells, called centroacinar cells, can give rise to  $\beta$ -cells in adult fish and mice (Delaspre et al., 2015; Mameishvili et al., 2019). Moreover, in juvenile mice,  $\delta$ -cells initiate insulin production upon ablation of the  $\beta$ -cells; however, this cellular makeover ceases to operate in older animals. Defining the full spectrum of cellular dynamics underlying  $\beta$ -cell regeneration requires technologies that can track gene expression in multiple cells over time during the process of  $\beta$ -cell regeneration.

In this study, we applied a unique blend of *in vivo* imaging, single-cell genomics and genetics to study *de novo*  $\beta$ -cell regeneration in zebrafish. Although regeneration is incomplete in mammals, zebrafish can naturally recover from extreme  $\beta$ -cell destruction and hyperglycemia (Moss et al., 2009). However, the reasons underlying this ability remain unexplored. By categorizing the cells in the zebrafish pancreas, we identify two discrete populations of  $\delta$ -cells, which express different *sst* paralogues. Moreover, we find that a subset of islet cells exists under a hybrid identity, sharing the hormones and fate determinants of both  $\delta$ - and  $\beta$ -cells. During ablation and subsequent regeneration, the hybrid cells form *de novo*, acquire glucose responsiveness and serve as a major source of insulin expression. We also show that the secreted protein Dkk3b can increase the formation of hybrid cells in the

absence of injury. We propose that the rapid formation of glucoseresponsive hybrid cells provides a shortcut to restoring glucose homeostasis and contributes to the ability of zebrafish to reverse the course of diabetes.

#### **RESULTS**

# Single-cell RNA-sequencing of the pancreas identifies two populations of $\delta\text{-cells}$ in zebrafish

We first characterized the cell types in the zebrafish pancreas under homeostasis. We analyzed the data from a droplet based single-cell transcriptome profiling of the islets and surrounding exocrine tissues from 2 months post-fertilization (mpf) zebrafish (Salem et al., 2019). For this experiment, the tissue surrounding the principal islet was dissected from the pancreas (n=6 animals). The dissected tissue was enzymatically dissociated, sorted by flow cytometry to remove damaged cells and profiled using the 10× Genomics pipeline (Fig. 1A). The transcriptome data was subjected to quality control that included a number of unique molecular identifiers (UMIs), genes detected per cell and the percentage of mitochondrial reads detected per cell. This yielded a total of 1969 cells for downstream analysis. Clustering of the single-cell transcriptome data using Seurat identified eight major clusters in the data (Fig. 1B-D). Based on the expression of marker genes, the clusters corresponded to acinar (ela2), ductal (cftr, hnflba, her15.1 and anxa4; Fig. S1A), endothelial (vsg1, also known as plvapb),  $\beta$ -cells (ins),  $\alpha$ -cells (gcgb),  $\varepsilon$ -cells (ghrl), and two discrete types of  $\delta$ -cells (sst1.1 and sst2) (Fig. 1B,C) (Table S1). The two groups of  $\delta$ -cells include a population marked by sst2 ( $\delta$ 2) (Li et al., 2009) and a distinct type that expresses sst1.1 ( $\delta$ 1). Each of the two sst genes, which arose through gene duplication in teleost fish, is predicted to encode for a full-length and functional somatostatin protein (Devos et al., 2002; Liu et al., 2010). One notable difference between the  $\delta 1$ - and  $\delta 2$ -cell types is the preferential expression of pancreatic and duodenal homeobox 1 (pdx1) in the former group (Fig. 2A). A number of other genes were differentially expressed between the two  $\delta$ -cell populations (Fig. 2B and Table S2). Thus, the zebrafish islet is equipped with two distinct δ-cell types, each expressing a different *sst* paralogue.

# The zebrafish islets contain a persistent bi-hormonal cell population

Most cell populations we identified contained discrete hormonal cell types. One exception occurs in the cells of the *ins* and *sst1.1* lineages. Notably,  $\sim$ 8% of all *ins*-expressing cells (60 cells out of 746) were positive for both *ins* and *sst1.1* (Fig. 1E). The bi-hormonal cells had further distinguishing features. The cells expressed *dkk3b*, encoding a Dickkopf-related secreted protein, and *pyyb*, a short peptide related to protein YY (Fig. 1E). *dkk3b* and *pyyb* were also enriched in  $\delta$ 1-cells compared to  $\delta$ 2-cells (Fig. 2).

To confirm the co-expression of *sst1.1* and *ins*, we used the single-cell transcriptomic profiles of sorted β-cells from seven stages of zebrafish life, including the juvenile and adult periods (Singh et al., 2018). We detected above-background expression of *sst1.1* in a fraction of *ins*-expressing cells at each of the seven stages (Fig. S2A). The fraction of *sst1.1*-expressing cells ranged from 5 to 9% of the *ins*-positive population throughout the life of the fish, showing that the bi-hormonal population maintains steady proportions.

Using assay for transposase-accessible chromatin (ATAC) of sorted  $\beta$ -cells from 2 mpf zebrafish, we detected open chromatin peaks in regulatory elements of *ins*, *sst1.1* and *dkk3b* (Fig. S2B). This suggests that *ins*-expressing cells have transcriptionally

favorable chromatin for expression of genes enriched in the bi-hormonal cells.

## Histological assessment of the bi-hormonal cells in the zebrafish pancreas

To directly visualize the bi-hormonal cells in the zebrafish pancreas, we took advantage of a transgenic reporter line, Tg(sst1.1: EGFP-Ras) (Löhr et al., 2018), in which the regulatory sequences of the sst1.1 gene drive a membrane-localized green fluorescent protein (Fig. 3A). Within islets, the GFP-positive  $\delta$ 1-cells formed an intricate mesh of cells distributed throughout the zebrafish islet and are interconnected by long cellular processes (Fig. 3B and Fig. S3A). We stained the pancreas from Tg(sst1.1: EGFP-Ras) larvae with antibodies against insulin to mark the  $\beta$ -cells. Confocal imaging of the islets demonstrated the presence of individual cells that co-expressed both EGFP and insulin (Fig. 3B). We further confirmed the co-expression of somatostatin and insulin proteins using immunofluorescence in pancreatic sections from adult fish aged 7 and 16 mpf (Fig. 3C,D). We clearly detected immunoreactivity for the two hormones along with the membrane GFP maker in single cells, corroborating the presence of bihormonal cells in the islet. In the bi-hormonal cells, insulin showed higher concentration in the cytoplasm, whereas somatostatin was localized closer to the cell membrane, which may reflect differential distribution of secretary granules for the two hormones. To investigate whether the ins/sst1.1 bi-hormonal cells occupy a specific position in the islet, we determined their distribution in adult zebrafish islets. We observed that the bi-hormonal cells are predominantly present in the periphery of the islet, with fewer cells in the center (Fig. S3B). Based on these results, along with the single-cell transcriptomic data, we have identified cells that express at the RNA and protein levels both insulin and somatostatin, and that persist from the early stages of development to adulthood.

# The bi-hormonal cells exhibit glucose-stimulated calcium influx in vivo

In order to investigate whether the bi-hormonal cells are glucose responsive and whether they present influx of calcium ions upon glucose stimulation, we performed 2-photon calcium imaging *in vivo* using Tg(sst1.1:EGFP-Ras); Tg(ins:K-GECO1) double transgenic larvae. We used our previously developed method for calcium imaging (Salem et al., 2019). We found that cells co-expressing EGFP and K-GECO1 show glucose-stimulated calcium influx, which was comparable with the response of  $\beta$ -cells in terms of speed and amplitude of response (n=3 islets) (Fig. 3E). Thus, the hybrid cells are functionally similar to  $\beta$ -cells in their capacity to respond to glucose.

# The bi-hormonal cells are molecular hybrids between $\delta \text{1-}$ and $\beta \text{-}\text{cells}$

To characterize the bi-hormonal cells at a better molecular resolution than the one provided by the  $10\times$  Genomics pipeline, we collected pancreatic tissue from 2 mpf fish and separated  $\beta$ -,  $\delta$ 1- and hybrid cells using index flow cytometry based on the expression of *ins*:tdTomato and *sst1.1*:EGFP-Ras (Fig. 4A). We performed single-cell transcriptomics using the Smart-Seq2 protocol, which is more sensitive than droplet-based single-cell sequencing (Ziegenhain et al., 2017). The single-cell sequencing revealed enrichment of *sst1.1* and *ins* in  $\delta$ 1-cells and  $\beta$ -cells, respectively, validating the utility of the reporter lines to distinguish the two cell populations (Fig. 4B,C). Furthermore, we observed a strong Pearson correlation in the co-expression of *sst1.1* and *dkk3b*, as well as of

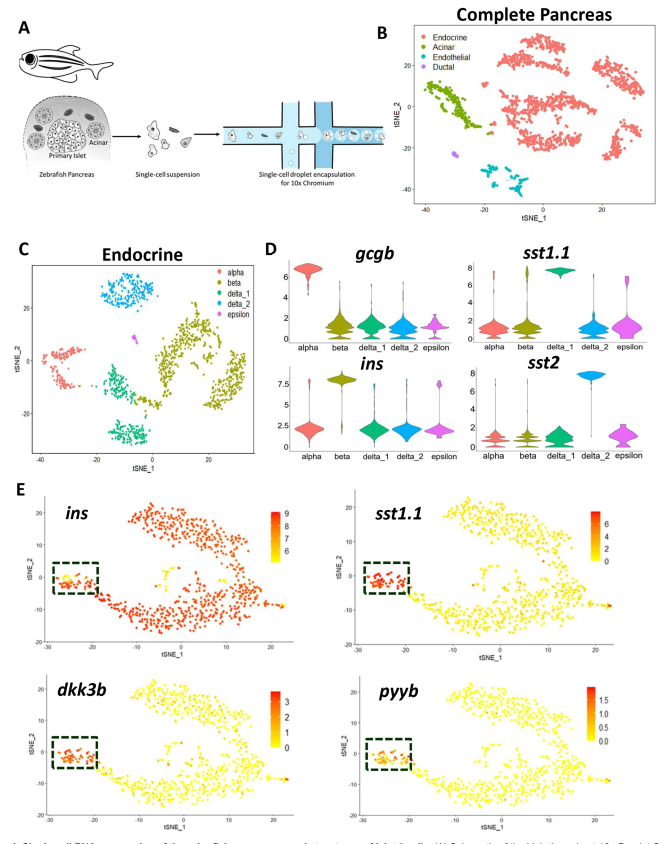
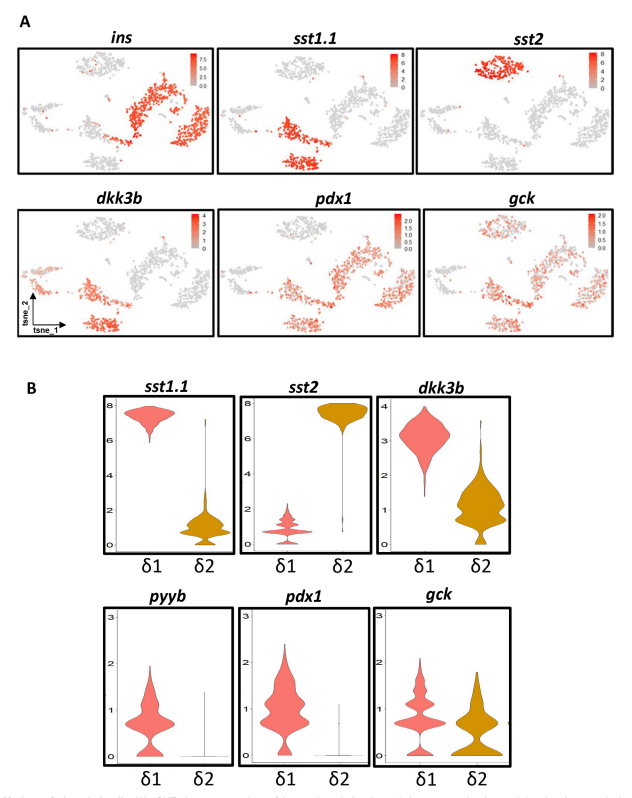


Fig. 1. Single-cell RNA-sequencing of the zebrafish pancreas reveals two types of islet  $\delta$ -cells. (A) Schematic of the high-throughput 10× Droplet-Seq approach to classify the cell populations in the pancreas from 2 mpf zebrafish. (B) t-SNE plot representation of 1969 pancreatic cells with clusters representing distinct cell types. (C) t-SNE plot representation of 1530 endocrine cells with clusters based on the expression of hormonal genes. (D) Violin plot for expression of hormonal genes in the endocrine clusters. (E) t-SNE plot representations of 746 insulin-expressing pancreatic cells with specific gene expression overlaid as a heat map. Cells with co-expression of *ins* and *sst1.1* are outlined.



**Fig. 2. Markers of δ1- and δ2-cells.** (A) t-SNE plot representations of the  $\beta$ -,  $\delta$ 1- and  $\delta$ 2-cell populations, expressing *ins*, sst1.1 and sst2, respectively. Glucokinase (gck) is expressed in both  $\delta$ 1- and  $\delta$ 2-cells, whereas dkk3b and pdx1 are enriched in the  $\delta$ 1-cells. (B) Violin plots highlighting the expression levels of sst1.1, sst2, dkk3b, pyyb, pdx1 and gck in the  $\delta$ 1- and  $\delta$ 2-cell clusters.

sst1.1 and pyyb in individual cells (Fig. 4D), validating our previous observations. Differential gene expression analysis showed that β-cells express significantly higher ( $P_{\rm adj}$ <0.05) levels of nkx6.2 and mnx1 (Fig. 4E), genes involved in their specification and maturation (Binot et al., 2010; Pan et al., 2015). Conversely, δ1-cells expressed

hhex (Fig. 4E), which encodes a transcription factor involved in the maintenance of δ-cell identity (Zhang et al., 2014). Interestingly, we observed intermediate expression levels for nkx6.2, mnx1 and hhex in the bi-hormonal cells, revealing an overlap of β- and δ-cell identity genes within the mixed population.

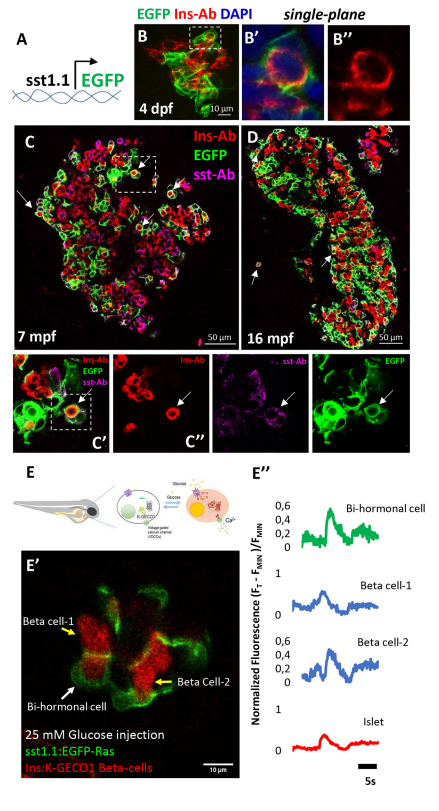


Fig. 3. Characterization of a small population of glucose-responsive β/δ1 hybrid cells. (A) Schematic of the Tg(sst1.1:EGFP-Ras) reporter line used to mark specifically the sst1.1 population, where a BAC containing the sst1.1 promoter drives membrane GFP expression. (B) Confocal projection of whole-mounted islets from the Tg(sst1.1: EGFP-Ras) reporter line stained with an antiinsulin antibody. The rectangle outlines a GFP and insulin co-expressing cell. The high-magnification images in B' and B" show a single confocal plane of the same cell coexpressing GFP and insulin. (C,D) Thin pancreatic sections from the Tg(sst1.1: EGFP-Ras) reporter line at 7 months post-fertilization (mpf, C) and 16 mpf (D). The sections were stained using insulin (red) and somatostatin (magenta) antibodies. Arrows indicate EGFP+ cells co-stained using insulin and somatostatin antibody. The rectangle in C outlines the region shown at higher magnification in C' and C", revealing cells expressing each protein along with GFP. (E-E") Two-photon in vivo Ca2+ imaging in 5 dpf doubletransgenic zebrafish larvae of the genotype *Tg(sst1.1*: EGFP-Ras); Tg(ins: K-GECO1), showing the glucosestimulated response of mono-hormonal and bi-hormonal cells (E'). Normalized K-GECO fluorescent traces showing glucose-stimulated calcium influx of bi-hormonal and monohormonal β-cells (E").

#### The bi-hormonal cells express endocrine-progenitor markers

Next, we applied pseudo-temporal analysis using Monocle (Trapnell et al., 2014) to model cellular transitions. The pseudo-temporal model showed that the bi-hormonal cells were positioned in between  $\beta$ - and  $\delta$ 1-cells (Fig. 4F), confirming the hybrid signature of these cells. We further characterized gene expression dynamics by calculating expression changes along the pseudotime

(Fig. 4F'). As expected, transcription factors implicated in  $\beta$ -cell lineage and identity (nkx6.2 and mnx1) were enriched towards the  $\beta$ -side of the pseudotime. Moreover, proprotein convertase subtilisin/kexin type 1 (pcsk1) (Benjannet et al., 1991), a gene necessary for conversion of pre-proinsulin to functional insulin hormone displayed a similar trend. In contrast, genes implicated in  $\delta$ -cell identity (hhex), along with pathways controlling endocrine

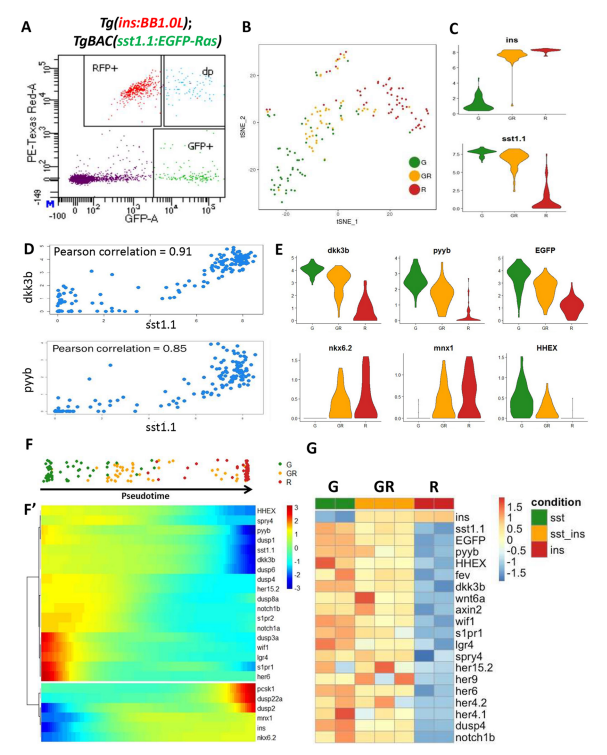


Fig. 4. Molecular analysis of the hybrid cells reveals a progenitor signature. (A) FACS plot of cells from Tg(ins:BB1.0L); Tg(sst1.1:EGFP-Ras) adult fish. Cells were marked based on the presence of red and green fluorescence, and index sorted into 96-well plates. β-Cells are RFP+ (R), δ1-cells are GFP+ (G) and bi-hormonal cells are green+ and red+ (GR). (B) A t-sne plot of the profiled cells with colors corresponding to their respective location on the FACS plot. The cells are segregated into three groups: R (β-cells), G (δ1-cells) and GR (bi-hormonal cells). (C) Violin plots showing *ins* and sst1.1 expression in the three cell populations. The *y*-axis represents normalized gene expression. (D) A dot-plot displaying the Pearson correlation coefficient between sst1.1 and dkk3b, as well as between sst1.1 and pyyb expression levels. (E) Violin plot showing the expression levels of control genes (dkk3b, pyyb and EGFP) and selected genes related to β-cell differentiation and function in the three populations. mnx1 and nkx6.2 are enriched in β-cells, whereas nhex is expressed in δ1-cells. The y-axis represents normalized gene expression. (F,F') Pseudo-temporal ordering of the cells (F); the expression trend of selected genes that show significant correlation with the pseudo-time (F'). The δ1-cells show enrichment of genes involved in the Wnt, FGF, Notch and sphingolipid signaling pathways compared with β-cells. The bi-hormonal cells exhibit intermediate expression of genes involved in β- and δ-cell fate specification and function compared with the pure cell populations. (G) To validate the single-cell RNA-Seq results, the three endocrine cell populations were profiled using bulk-RNA sequencing. The results from the experiment are depicted as a heat-map demonstrating the expression levels of selected genes, including those involved in Wnt, FGF, Notch and sphingolipid signaling. All depicted genes show a statistically significant difference in expression between the  $\delta$ 1- and  $\beta$ -cells.

progenitor maintenance, such as Wnt, Fgf, Notch and sphingolipid signaling (sIpr1 and sIpr2) were associated with the trajectory towards the δ1-cells (Ninov et al., 2012; Serafimidis et al., 2017; Seymour et al., 2012; Sharon et al., 2019; Zhang et al., 2014). The hybrid cells expressed intermediate levels of pathway components related to these pathways, such as dusp3a, wif1, lgr4, sIpr1 and her6, when compared with either β- or δ1-cells. The expression of the pathway components involved in progenitor maintenance was confirmed by bulk RNA-sequencing of the sorted cells (Fig. 4G), which also revealed enrichment of fev, a transcription factor that was recently shown to be expressed in endocrine progenitors in mice (Byrnes et al., 2018). Altogether, we conclude that the hybrid cells display the transcriptional profiles of intermediate cell states between δ1- and β-cells, and exhibit an enrichment of pathways associated with endocrine progenitors.

#### The hybrid cells increase during $\beta$ -cell regeneration

Intrigued by the association of precursor characteristics of hybrid cells, we further characterized their behaviors during β-cell regeneration in larval zebrafish. Near-complete β-cell death was induced using the nitroreductase (NTR) system (Curado et al., 2007; Pisharath et al., 2007). In this system, Tg(ins:FLAG-NTR) transgenic zebrafish, expressing NTR specifically in the β-cells, are exposed to Mtz (metronidazole). NTR converts Mtz to a cytotoxin, which specifically kills the  $\beta$ -cells while sparing all other cells in the body. To label sst1.1-expressing cells during  $\beta$ -cell regeneration, we used a Tg(ins:FLAG-NTR); Tg(sst1.1:EGFP-Ras) double transgenic line. We exposed Tg(ins:FLAG-NTR) larvae to Mtz for 24 h, and collected samples for observation 4 and 8 days post ablation (dpa) (Fig. S4A). At both stages, we observed a dramatic increase in the percentage of hybrid cells within the regenerating islet (Fig. S4B,C). Of note, when we examined the expression of sst2:RFP, which marks the population of  $\delta$ 2-cells ( $\delta$ 2) in zebrafish (Li et al., 2009), we did not observe the formation of hybrid cells (O.K., unpublished).

# A single-cell map of adult $\beta$ -cell regeneration reveals a major contribution of hybrid cells to insulin expression

To assess the dynamics of hybrid cells during the more physiologically-relevant setting of adult β-cell regeneration, we performed single-cell RNA sequencing (using the 10× genomics pipeline) at different times following β-cell destruction and regeneration in adult zebrafish (9 mpf of age). We selectively ablated the β-cells and dissected the primary islets and surrounding tissues, including the exocrine cells from adult zebrafish at 0, 2, 7 and 14 dpa. These time points were chosen as they include the time of β-cell destruction, subsequent hyperglycemia, followed by normalization of glucose levels after 2 weeks of regeneration (Fig. S5) (Delaspre et al., 2015). To induce efficient ablation of insulin-expressing cells, we used a new ablation line, which employs an improved and more effective version of the NTR enzyme (Fig. S5) (Singh et al., 2017). In the unablated samples, we identified all major cell types, including discrete populations of *ins*-expressing  $\beta$ -cells, *sst1.1*-expressing  $\delta$ -cells, *sst2*-expressing δ2-cells, ductal and acinar cells (Fig. 5 and Table S3). Shortly upon β-cell ablation (at 0 and 2 dpa), the *ins*-only expressing population observed in controls became nearly completely absent, based on single-cell gene expression analysis. Instead, ins expression was detected mainly within the  $\delta 1$  population, in which cells coexpressed both ins and sst1.1 (Fig. 5 and Fig. S6). At 7 and 14 dpa, the hybrid cells increased further due to greater proportions of insexpressing cells in the cluster corresponding to  $\delta$ 1-cells (Fig. S7).

Clustering of the integrated data from all time-points revealed that the hybrid cells within the sst1.1 cluster are the main source of insulin expression during regeneration (Fig. 5D,E). Moreover, upon  $\beta$ -cell destruction, we detected increased expression in pcsk1 and pcsk2 in the sst1.1-expressing cells, which are necessary for insulin maturation (Fig. S8). These changes in gene expression were accompanied by an increase in the number of sst1.1-expressing cells relative to ablation-invariant  $\alpha$ -cells (Fig. S9). At 7 dpa, the relative number of sst1.1-expressing cells has stabilized at roughly twice as many as in controls.

Compared with the sst1.1-expressing cells,  $\delta$ 2- and  $\alpha$ -cells showed a relatively minor contribution to insulin expression (Fig. 5D,E). A closer inspection of the  $\alpha$ -cell cluster highlighted that some gcgbexpressing cells do show low-to-medium ins expression over the course of β-cell regeneration (Fig. S10A-D). This suggests that individual α-cells also act as sources of insulin expression. However, compared with the  $\delta$ 1-cells,  $\alpha$ -cells are not the major contributors. Finally, we generated a list of differentially expressed genes (DEGs) in  $\alpha/\beta$  and  $\delta/\beta$  hybrid cells at 7 dpa and performed Gene Ontology (GO) analysis (Table S4). Whereas  $\delta/\beta$  hybrid cells showed an enrichment of genes involved in biological processes, including 'type B pancreatic cell differentiation', 'activation of MAPKKK activity' and 'negative regulation of canonical Wnt signaling', the α/β hybrid cells were associated with genes linked to 'fatty acid elongation', 'protein peptidyl-prolyl isomerization', 'positive regulation of cytokine production and angiogenesis' and 'neuropeptide signaling pathway'. This analysis hints that different molecular mechanisms underlie the formation of  $\alpha/\beta$  and  $\delta/\beta$  hybrid cells (Fig. S10E).

#### The hybrid cells retain bi-hormonal characteristics

To investigate whether  $\delta/\beta$  hybrid cells may resolve into sst1.1<sup>+</sup> and ins<sup>+</sup> populations during regeneration, we first analyzed the single-cell transcriptome data from 7 and 14 dpa (Fig. 6). 7 dpa represents an intermediate stage whereas 14 dpa corresponds to the resolution of diabetes. By specifically focusing on cells that express the two hormones, we observed overlap between the sst1.1<sup>+</sup> and ins<sup>+</sup> cells at 7 dpa (Fig. 6A,B). By contrast, at 14 dpa the cells segregated into two clusters (Fig. 6D,E). Notably, the two clusters observed at 14 dpa were enriched for the expression of either sst1.1 or ins (Fig. 6E), indicating that the two cell populations were diverging. We further performed gene correlation analysis to examine the distribution of sst1.1 and ins gene expression level at cellular resolution. At 7 dpa, a positive correlation was observed between ins and sst1.1 expression (Fig. 6C), suggesting the presence of hybrid cells at this stage. At 14 dpa, however, individual cells displayed a negative correlation between sst1.1 and ins expression (Fig. 6F), suggesting a trend towards the resolution of gene expression into  $\beta$ - or  $\delta$ 1-cells.

To corroborate our observations from the transcriptome analysis, we co-stained the adult zebrafish pancreatic tissues sections using somatostatin and insulin antibody. Compared with the control samples where cells distinctly expressed insulin and somatostatin, we observed a near to complete loss of insulin-positive cells at 2 dpa (Fig. S11A,B). At 7 and 14 dpa, we found that the insulin-expressing cells co-expressed somatostatin. Notably, the majority of the insulin-positive cells continued to exhibit bi-hormonal signature until 14 dpa (Fig. S11C,D). Thereafter, we also characterized the secondary islets over the course of regeneration. Similar to the primary islets, the secondary islets in the unablated samples did not show a significant overlap between the somatostatin and insulin-positive cells. However, there was a predominance of ins/sst bi-hormonal cells both at 7 dpa and at 14 dpa (Fig. S12). Thus, the immunohistological results contrast with the transcriptomics

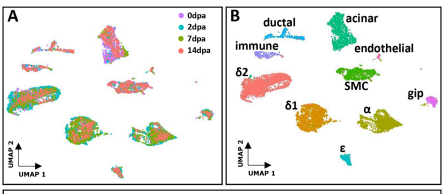
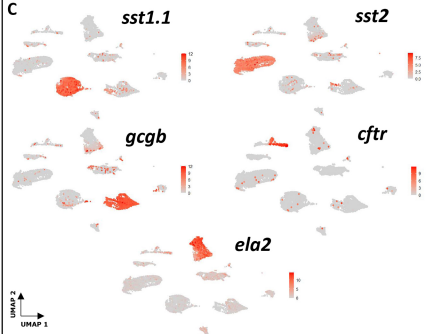
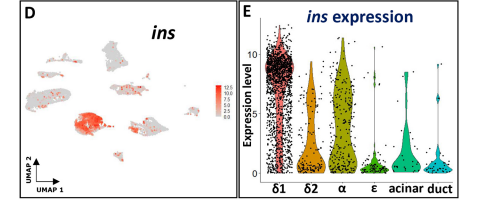


Fig. 5. The hybrid cells constitute a prominent source of insulin expression during diabetes recovery in adult zebrafish. (A,B) UMAP plot representation of integrated 10× scRNA-Seq data from all  $\beta$ -cell regeneration time points 0, 2, 7 and 14 days post-ablation (dpa) showing distinct clusters for  $\delta$ 1,  $\delta$ 2,  $\alpha$ , acinar and duct cells. (C) Feature-plot highlighting the expression of sst1.1 ( $\delta$ 1), sst2 ( $\delta$ 2), gcgb ( $\alpha$ ), cftr (duct) and ela2 (acinar) genes. (D) Feature-plot highlighting ins expression in different pancreatic cells during regeneration (E) Violin plot highlighting ins expression in different pancreatic cells. The  $\beta$ / $\delta$ 1 hybrid cells among the  $\delta$ 1-cells serve as the main source of insulin expression during the course of regeneration.





analysis and indicate that the resolution of the cells into monohormonal cells is incomplete until at least 2 weeks post-β-cell loss.

# Live-cell tracking reveals hybrid-cell transition during regeneration

To assess if the bi-hormonal cells arise de novo during  $\beta$ -cell regeneration or whether they might reflect pre-existing cells that have survived the ablation procedure, we performed live imaging of islets undergoing regeneration during the optically translucent larval stage. Insulin-expressing cells were labeled using ins:tdTomato, whereas

sst1.1-expressing cells were marked using sst1.1:EGFP-Ras. Ablation of β-cells was induced using Tg(ins:FLAG-NTR) by Mtz-treatment from 3 to 4 dpf. First, we measured the efficiency of cell ablation and found that both β-cells and hybrid cells were efficiently ablated (Fig. 7A). We then monitored longitudinally the regeneration process in individual larvae. Strikingly, δ1-cells showed dramatic changes in morphology, as they accumulated prominent membrane-bound vacuolar structures within their cytosol, evident both under confocal and electron microscopy (Fig. S13). Moreover, within islets that previously had lost any tdTomato expression, we now observed

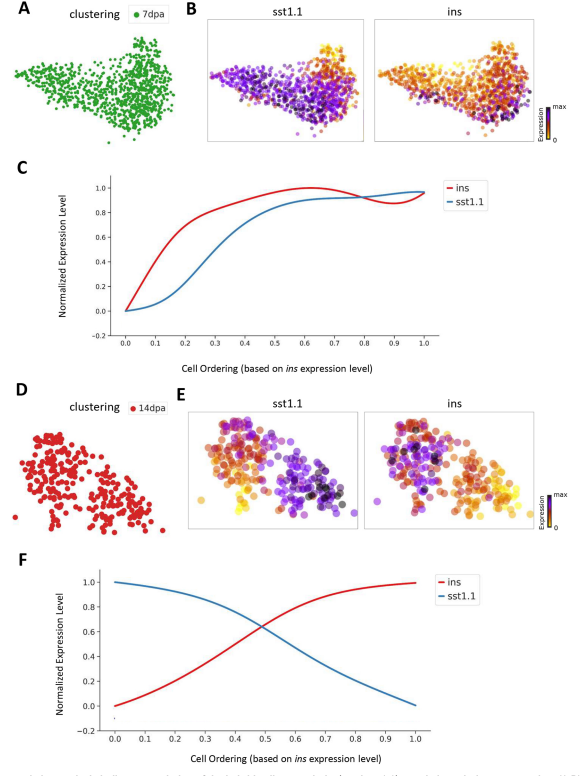


Fig. 6. Gene correlation analysis indicates resolution of the hybrid cells towards *ins*<sup>+</sup> and *sst1.1*<sup>+</sup> populations during regeneration. (A,D) UMAP displaying selected cells from 7 dpa (A) and 14 dpa (D). (B,E) UMAP overlaid with gene expression plots for *sst1.1* and *ins* at 7 dpa (B) and 14 dpa (E). (C,F) Expression trends for *sst1.1* and *ins* at 7 dpa (C) and 14 dpa (F). At 14 dpa, individual cells along display a negative correlation between *sst1.1* and *ins* expression, whereas at 7 dpa the cells co-express both hormones.

its re-expression in EGFP-positive cells, indicating the *de novo* origin of the hybrid cells (Fig. 7A).

Furthermore, we wanted to assess whether pre-existing  $\beta$ -cells, spared from ablation, may undergo de-differentiation to form

Ins/Sst bi-hormonal cells. To this end, we performed  $\beta$ -cell tracing using Tg(ins: H2B-mEos2); Tg(ins: FLAG-NTR) double-transgenic zebrafish larvae. These larvae express a green-to-red photoconvertible protein mEos2 (fused to histone H2B) and NTR

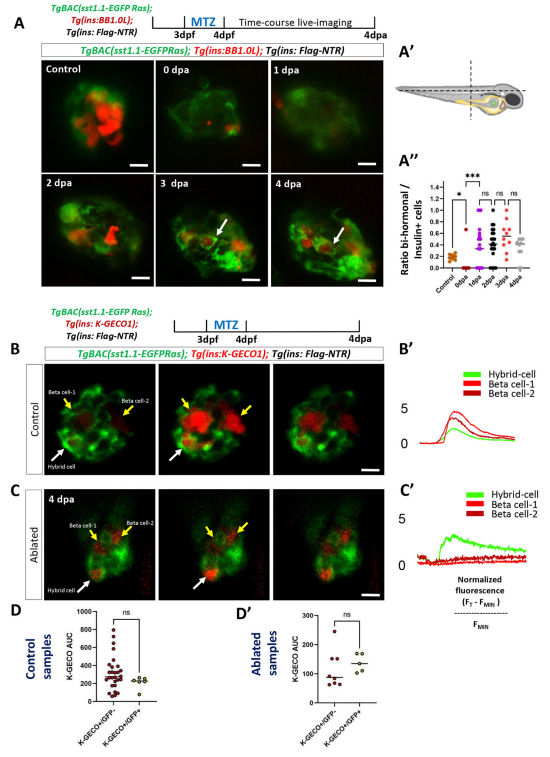


Fig. 7. Live imaging of regeneration reveals that the hybrid cells arise *de novo* and acquire glucose responsiveness. (A) To visualize the dynamics of bi-hormonal cells during β-cell regeneration, Tg(ins: FLAG-NTR); Tg(sst1.1: EGFP-Ras); Tg(ins: BB1.0L) triple transgenic animals (5 dpf) were treated with vehicle or Mtz for 24 h. Following Mtz-mediated ablation of β-cells, a time-course of live imaging was performed from 0 to 4 dpa. 3D projections of the zebrafish islet in control and at different time-points post β-cell ablation in the same larva. Following Mtz addition, all tdTomato-expressing cells were ablated and no longer present in the islet. New tdTomato- and GFP-positive cell appear in the islet over time, indicating a *de novo* origin (arrows). (A') Orientation of the zebrafish islet during the live-imaging process. (A") Quantification of the proportion of bi-hormonal cells during the regeneration process (\*P<0.05, \*\*\*P<0.001; Brown-Forsythe and Welch ANOVA). (B,C) Images from the time-lapse recording of glucose-stimulated calcium influx (six frames/s, single plane) of the zebrafish islet in control (unablated) and at 4 days post β-cell ablation (ablated). (B',C') Normalized *K-GECO* fluorescent traces showing glucose-stimulated calcium influx of bi-hormonal and monohormonal cells. (D,D') Quantification of changes in normalized *K-GECO* fluorescence intensity (recorded over 400 frames after glucose injection) based on area under the curve (AUC) for single cells. (D) Scatterplot for *K-GECO+/GFP*<sup>-</sup> and *K-GECO+/GFP*<sup>+</sup> from unablated samples (Mann–Whitney unpaired *t*-test, *P*=0.0907, non-significant). Scatterplot for *K-GECO+/GFP*<sup>-</sup> and *K-GECO+/GFP*<sup>+</sup> from ablated samples (Mann–Whitney unpaired *t*-test, *P*=0.222, non-significant). Scate bars: 5 μm.

specifically in the  $\beta$ -cells. At 3 dpf, we photolabeled all pre-existing  $\beta$ -cells in red, followed by Mtz treatment. At 3 dpa, the majority of the hybrid cells displayed green nuclei and somatostatin expression. This strongly suggests that the bi-hormonal cells do not arise from the pre-existing  $\beta$ -cells in the zebrafish larvae (Fig. S14).

## The newly emerging bi-hormonal cells exhibit glucose-stimulated calcium influx in vivo

In order to investigate whether the emerging bi-hormonal cells, which arise post β-cell ablation, are glucose responsive and present a calcium influx upon glucose stimulation, we performed in vivo calcium imaging using Tg(sst1.1:EGFP-Ras); Tg(ins:FLAG-NTR) and Tg(ins:K-GECO1) triple-transgenic larvae. We specifically ablated  $\beta$ -cells by exposing the larvae to Mtz from 3 to 4 dpf and performed in vivo calcium imaging at 2, 4 and 6 days post β-cell ablation (Movies 1 and 2). Interestingly, we found that the emerging bi-hormonal cells became glucose responsive as early as 4 days post-β cell ablation (Fig. 7B-C) (Movie 2). The bi-hormonal cells showed a comparable glucose responsiveness to GFP-negative but K-GECO1-positive cells (Fig. 7D), which likely arise via constitutive developmental neogenesis in larvae. The glucose responsiveness of bi-hormonal cells, however, was lower than that of β-cells in unablated animals, likely because the cells were undergoing maturation. Thus, our results indicated that the emerging bi-hormonal cells acquire responsiveness to glucose stimulus.

#### Overexpression of dkk3b increases hybrid-cell formation

Our initial analysis of gene expression of the hybrid cells indicated that the cells are enriched for the expression of dkk3b, which is also shared with  $\delta 1$ -cells. Moreover, upon  $\beta$ -cell ablation, dkk3b was among the genes that showed a trend of increasing expression within the  $\delta 1$ -cell cluster over time (Fig. S8). To test whether dkk3b plays a role in hybrid-cell formation, we generated transgenic lines with constitutive expression of dkk3b under the ins promoter, or with inducible expression based on the heat-shock promoter (hsp). In both cases, we observed that the upregulated expression of dkk3b in the islets let to increased proportions of insulin and somatostatin double-positive cells during development (Fig. 8A,B). Thus, the forced overexpression of dkk3b, which is normally enriched in hybrid and  $\delta 1$ -cells, was sufficient to promote the formation of bi-hormonal cells without injury.

#### Zebrafish $\delta$ 1-cells are partially related to $\gamma$ -cells in mammals

To identify the cellular counterparts of zebrafish  $\delta 1$ - and  $\delta 2$ -cells in the mammalian pancreas, we compared our single-cell transcriptomic dataset (Salem et al., 2019) with human pancreatic data (Segerstolpe et al., 2016). Based on the expression of marker genes in the human dataset, the cells were clustered into  $\alpha$ -cells (GCG),  $\beta$ -cells (INS),  $\delta$ -cells (SST),  $\epsilon$ -cells (GHRL),  $\gamma$ -cells (PYY), ductal (CFTR) and acinar (PRSS1) cells. The interspecies comparison revealed that zebrafish pancreatic cells clustered with their respective human counterparts, highlighting transcriptomic similarity (Fig. 9A). Notably, we observed that zebrafish  $\delta$ 2-cells clustered near to human  $\delta$ -cells, whereas the  $\delta$ 1-cells showed partial overlap with human y-cells. Moreover, there was a higher number of co-expressed genes between zebrafish δ2-cells and human  $\delta$ -cells, when compared with  $\delta$ 1-cells (Fig. 9B and Table S5). Finally, zebrafish  $\delta 1$ - but not  $\delta 2$ -cells express *pvvb*. encoding peptide YY, a peptide that is also expressed in mouse γ-cells (Perez-Frances et al., 2021) (Table S2 and Discussion). This analysis suggests that  $\delta$ 1-cells share transcriptional similarity to γ-cells in mammals.

#### **DISCUSSION**

In this study, we used single-cell sequencing to characterize cellular dynamics in a regeneration-competent animal model – the zebrafish (Fig. 8C). Using this approach, we defined the gene expression of different pancreatic cells upon  $\beta$ -cell destruction and established an atlas of  $\beta$ -cell regeneration, spanning the period from  $\beta$ -cell destruction to emergence of new insulin-expressing cells. Among the different cell types, we characterized a novel population of  $\delta$ -cells, expressing the *sst* paralogue *sst1.1*, which is consistent with previous studies showing its expression in the zebrafish pancreas (Dalgin et al., 2011; Devos et al., 2002; Spanjaard et al., 2018). In addition, the single-cell resolution of our analysis allowed us to pinpoint a small population of bi-hormonal cells, which exist under a hybrid  $\beta/\delta 1$ -cell identity.

#### Hybrid cells as an intermediate hormonal cell population

We show that the hybrid cells share the hormones, fate determinants and glucose responsiveness of both  $\delta 1$ - and  $\beta$ -cells. Although the cells are transcriptionally distinct from β-cells, they show glucosestimulated Ca<sup>2+</sup> influx. We validated the presence of hybrid cells at different stages of zebrafish life: from 4 dpf to 16 mpf. Moreover, using single-cell RNA-sequencing, we compared the transcriptional profile of the hybrid cells to mono-hormonal  $\beta$ -cells and  $\delta$ 1-cells. The differential gene expression analysis revealed an overlapping expression of genes involved in both  $\beta$  and  $\delta$ -cell fate-specification and function. For example, the cells exhibit intermediate expression of genes involved in insulin prohormone processing, when compared with β-cells. Moreover, these cells express at intermediate levels transcription factors that differentiate between the  $\beta$ - and  $\delta$ -cell lineages, such as the evolutionary conserved homeobox mnx1 and hhex, respectively (Dalgin et al., 2011; Pan et al., 2015; Zhang et al., 2014). In parallel work aimed to define the lineage decisions in the developing endoderm, we find that the population of sst1.1/ins cells becomes evident as early as the embryonic islet forms, maintaining similar proportions as in the mature adult islet (~8% of the insulin-expressing cells) (M.K., unpublished). Moreover, both sst1.1 and dkk3b genes were enriched in the β-cell precursor cluster in a previously published dataset from the embryonic pancreas (Lu et al., 2019). Together, the deep-sequencing experiments suggest that the hybrid cells are an intermediatelineage population.

#### Hybrid cells in rapid resolution of diabetes in zebrafish

Polyhormonal endocrine cells have been observed during embryonic pancreas development in mammals and during regeneration (De Krijger et al., 1992; Teitelman et al., 1993). However, it remains unknown whether such cells play a significant role in the pancreas. To test their role, we performed two parallel experiments. First, we asked whether the hybrid cells are glucose responsive. We found that under steady state, the cells behave similar to  $\beta$ -cells, as they exhibit a robust glucose-stimulated calcium influx. Moreover, during regeneration, the emerging hybrid cells acquire the capacity to undergo calcium influx in response to glucose stimulation. The functional role of the cells in the context of regeneration is supported by the second experiment we carried out: to follow the time course of regeneration using single-cell transcriptomics and to correlate insulin expression with reversal of diabetes. Although there was no evidence for the presence of a major group of regenerated mono-hormonal β-cells even 14 days after  $\beta$ -cell destruction, by this time, the fasting glucose was back to normal, with insulin being expressed predominately in the hybrid cells. Moreover, there was increased expression of pdx1 and pcsk1, which is in line with them acting to restore the insulin-

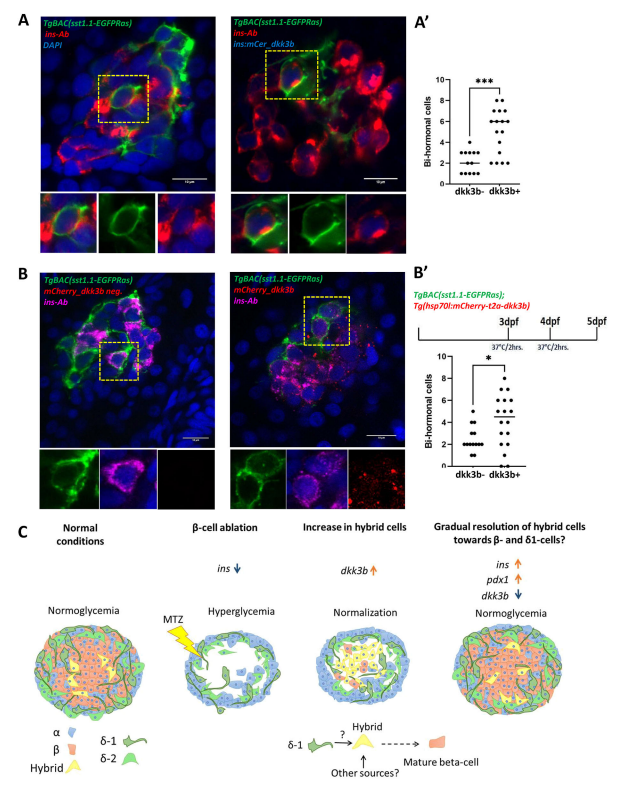


Fig. 8. Overexpression of *dkk3b* leads to increased formation of hybrid cells. (A) Single-plane confocal images of islets from *Tg(ins: mCerulean\_dkk3b)*; *Tg(sst1.1: EGFP-Ras)* double transgenic animals (5 dpf) stained using insulin antibody (red). The yellow dashed rectangle outlines the *sst1.1-ins* double-positive cells (shown below). (A') Scatter plot showing the number of bi-hormonal cells in wild-type and *ins:mCerulean-t2a-dkk3b* animals (Mann–Whitney unpaired *t*-test, \*\*\*\*P=0.0005, significant). (B) Single-plane confocal images of islets from *Tg(hsp70l:mCherry-t2a-dkk3b)*; *Tg(sst1.1: EGFP- Ras)* double transgenic animals stained using an insulin antibody (magenta). The larvae were given a heat-shock for 2 h at 37°C at 3 dpf and 4 dpf, thereby inducing the expression of *dkk3b*. The yellow dashed rectangle outlines the *sst1.1-ins* double-positive cells (shown below). (B') Scatter plot showing the number of bi-hormonal cells in wild-type and *Tg(hsp70l:mCherry-t2a-dkk3b)* larvae (Mann–Whitney unpaired *t*-test, *P*=0.0352, significant). (C) Cartoon model based on the experimental observations in this study. Scale bars: 10 μm.

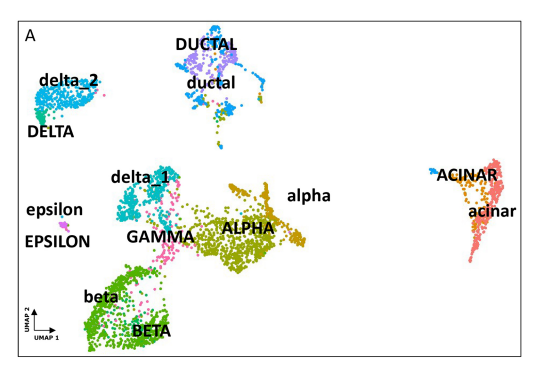


Fig. 9.  $\delta$ 1-cells are partially related to human  $\gamma$ -cells. (A) UMAP plot representation of integrated human and zebrafish datasets. Uppercase indicates human cell types and lowercase indicates zebrafish cell types. (B) Table highlighting the number of co-expressed genes for zebrafish and human cell types.

В	Cell Types	Co-expressed genes
	delta_1 vs DELTA	252
	delta_2 vs DELTA	353
	delta_1 vs GAMMA	319
	delta 2 vs GAMMA	304

producing function of the islet. Therefore, we show that the hybrid cells fulfill a role in glucose regulation in the absence of  $\beta$ -cells.

#### Hybrid cells inform on islet-cell plasticity

An important unanswered question remains as to the cellular sources of the hybrid cells during regeneration. Our observations point towards the subpopulation of sst1.1-expressing cells, which we suspect may increase insulin production upon  $\beta$ -cell destruction, thereby forming hybrid cells. Intriguingly, in mice,  $\delta$ -cells have been shown to be plastic in juvenile animal but lose their plasticity in adulthood (Chera et al., 2014). It is possible that adult zebrafish maintain an active population of  $\delta$ 1-cells, which covert into hybrid cells and make it possible to restore normoglycemia. However, we cannot exclude the possibility that the centroacinar subpopulation of terminal duct cells also undergo a process of endocrine differentiation, passing through a bi-hormonal state, which may further resolve into  $\beta$ -cells over time. Differentiating between these two sources will required the development of tools for lineage tracing of  $\delta$ 1-cells in the zebrafish pancreas.

A second important question will be to define the signaling pathways that promote the formation and resolution of such hybrid cells. Here, we show that hybrid cells express dkk3b, and its overexpression is sufficient to generate hybrid cells in the absence of injury. As dkk3b is predicted to be a secreted factor, the precise cell population that responds to its overexpression to generate hybrid cells remains to be defined. dkk3b was previously shown to be expressed in an unknown group of islet cells in zebrafish (Untergasser et al., 2011), and we now reveal its association with the sst1.1-expressing  $\delta$ -cells. Moreover, the human DKK3 is one of

the first examples of a heterogeneously expressed gene in islets (Hermann et al., 2007). Notably, *DKK3* is also upregulated in islets from donors with type 2 diabetes (Solimena et al., 2018). The role of this interesting gene in the context of diabetes remains to be studied.

#### Hybrid cells and the evolution of endocrine cell types

Our study also raises a curious question regarding the evolution of the islet cells types. Specifically, our interspecies comparison of gene expression illustrated that  $\delta$ 1-cells in zebrafish may be partially related to  $\gamma$ -cells. In mouse,  $\gamma$ -cells are marked by the expression of Pvv together with Ppv. PPY and PYY arose from the tandem geneduplication of the PYY gene (Perez-Frances et al., 2021; Hort et al., 1995). Similarly, we found that zebrafish  $\delta$ 1-cells also expressed the paralog pyyb. Thus, it is possible that the  $\delta$ 1-cells in zebrafish represent the evolutionary couterpart to γ-cells in mammals. A comprehensive review by Heller noted that the first examples of an islet-like organ containing three distinct endocrine cell types emerged in cartilaginous and bony fish. Later, amphibians and reptiles evolved islets with four major endocrine hormones, and in some but not all mammalian species, islets with all five cell types can be observed (Heller, 2010). We propose that the establishment of hybrid-cell populations may represent an intermediate stage along the diversification and evolution of the discrete subsets of endocrine cells.

Finally, one of the take-home messages is that regenerating imperfect yet functional hybrid cells capable of insulin-production may offer the opportunity to elicit glucose control. Perhaps, such surrogate cells would be stealthy in terms of escaping immune cell

recognition and destruction in type 1 diabetes, as they exist under a hidden identity, resembling two different hormonal cell types at the same time.

#### **MATERIALS AND METHODS**

#### Zebrafish strains and husbandry

Wild-type or transgenic zebrafish of the outbred AB, WIK or a hybrid WIK/AB strain were used in all experiments. Zebrafish were raised under standard conditions at 28°C. Animals were chosen at random for all experiments. Published transgenic strains used in this study were Tg(ins:BB1.0L; cryaa:RFP) (Singh et al., 2017),  $Tg(sst1.1:EGFP-Ras)^{fr40Tg}$  (Löhr et al., 2018),  $Tg(fabp10:dsRed, ela31:GFP)^{gz12}$  (Wan et al., 2006),  $Tg(ins:FLAG-NTR)^{s950}$  (Andersson et al., 2012),  $Tg(ins:YFP-2A-NTR3, cryaa: mCherry)^{tud201Tg}$  (Singh et al., 2017) and Tg(sst2:RFP) (Li et al., 2009). Experiments were conducted in accordance with the Animal Welfare Act and with permission of the Landesdirektion Sachsen, Germany (permits: TVV38/2015, TVV 45/2018, TVV33/2019, TVV 32/2020 and all corresponding amendments).

## Single-cell suspension of zebrafish pancreatic islets and liver cells

Single cell suspension of zebrafish pancreas was performed according to the protocol outlined previously (Singh et al., 2018). In brief, primary islets and substantial surround tissue were dissected and dissociated into single cells by incubation in TrypLE (Thermo Fisher Scientific, 12563029) with 0.1% Pluronic F-68 (Thermo Fisher Scientific, 24040032) at 37°C in a benchtop shaker set at 450 rpm for 30 min. During the dissection, we used the fluorescent signal of β-cells in the Tg(ins:BB1.0L; cryaa:RFP) or Tg(ins:YFP-2A-NTR3,cryaa:mCherry) reporters in order to orient ourselves with respect to the anatomical position of the main pancreatic islets. Following dissociation, TrypLE was inactivated with 10% FBS, and the cells pelleted by centrifugation at 500 g for 10 min at 4°C. The supernatant was carefully discarded and the pellet re-suspended in  $500\,\mu l$  of HBSS (without Ca, Mg) and 0.1% Pluronic F-68. To remove debris, the solution was passed over a 30 µm cell filter (Miltenyi Biotec, 130-041-407). To remove dead cells, calcein violet (Thermo Fisher Scientific, C34858) was added at a final concentration of 1 µM and the cell suspension incubated at room temperature for 20 min. The single cell preparation was sorted with the appropriate gates for identification of β-cells and alive cells (calcein<sup>+</sup>). FACS was performed through a 100 µm nozzle.

For samples from  $\beta$ -cell regeneration time-points, the adult fish were treated with Mtz (Sigma-Aldrich, M3761,) which was dissolved at 10 mM concentration in fish water. The animals were treated for 24 h, protected from light. After 24 h, Mtz solution was removed and the animals were placed three times in fresh fish water before returning them to standard maintenance conditions. Both untreated and Mtz-treated animals were euthanized by an overdose of tricaine. Samples for each time-point comprised six animals (three males and three females). Single-cell suspension was prepared in the same manner (as highlighted above) and 15,000 live cells were FACS sorted for  $10\times$  Genomics.

#### Single-cell RNA-Seq of the zebrafish pancreas

The sorted cells (38 µl) were carefully mixed with reverse transcription mix before loading the cells on the 10× Genomics Chromium system (Zheng et al., 2017) in a Chromium Single Cell B Chip and processed further following the guidelines of the 10× Genomics user manual. In short, the droplets were directly subjected to reverse transcription, the emulsion was broken and cDNA was purified using Silane beads. After the amplification of cDNA with 11 cycles, it underwent purification and quantification. The 10× Genomics single cell RNA-seq library preparation – involving fragmentation, dA-Tailing, adapter ligation and a 13 cycle indexing PCR – was performed based on the manufacturer's protocol. After quantification, the libraries were sequenced on an Illumina Novaseq6000 in pairedend mode (R1: 29 cycles; I1: 8 cycles; R2: 93 cycles), thus generating ~50-150 mio. fragments. The raw sequencing data was then processed with the 'count' command of the Cell Ranger software (v4.0.0) provided by 10× Genomics with the option '–expect-cells' set to 7500 (all other

options were used as per default). To build the reference for Cell Ranger, zebrafish genome (GRCz11) as well as gene annotation (Ensembl 95) were downloaded from Ensembl and the annotation was filtered with the 'mkgtf' command of Cell Ranger (options: '-attribute=gene\_biotype:protein\_coding -attribute=gene\_biotype: lincRNA -attribute=gene\_biotype:antisense'). Genome sequence and filtered annotation were then used as input to the 'mkref' command of Cell Ranger to build the appropriate Cellranger Reference.

#### Analysis of single-cell RNA-Seq of the zebrafish pancreas

We re-analyzed the single-cell RNA-Seq of the zebrafish pancreatic cells, which was generated using the 10× Chromium pipeline and previously published (Salem et al., 2019). The data have been deposited in GEO under accession number GSE123662. For clustering and gene expression analysis, we used Seurat and followed the recommended analysis pipeline. Briefly, the raw data as UMI counts was log normalized, regressed to remove the effect of library size and scaled. Highly variable genes were identified for PCA analysis and graph-based clustering. Marker genes identified for each cluster were used to classify the exocrine, endocrine and ductal population. The endocrine cluster was isolated and sub-clustered to identify and label individual endocrine populations. The cells were filtered using the following QC metrics: 3000<nFeature\_RNA>350, 25,000<nCount\_RNA>2500 and percent.mito<20. All the cells that passed the quality check were used for downstream analysis.

To define the changes the abundance of  $\delta 1$ -cells during regeneration, at each time point, we calculated the ratio between the number of  $\delta I$ - and  $\alpha$ -cells, filtering out any clusters that expressed multiple hormones apart from the combination of sst1.1 and ins. We calculated 99% confidence intervals from a 1000-fold random sampling of the total number of cells present at every time from a binomial distribution with the observed cell type frequencies as probabilities, using the function rbinom in base R.

#### Comparison of zebrafish and human datasets

To identify transcriptional overlap between cell types from human and zebrafish pancreas, we quantified the number of co-expressed genes between cell clusters from single-cell dataset of zebrafish (Salem et al., 2019) and human (Segerstolpe et al., 2016). We converted the human genes to their corresponding zebrafish homologues using the mapping from bioMart. For human genes with multiple zebrafish orthologues, the gene was mapped to each orthologue. Next, we integrated the modified human dataset and the zebrafish dataset using ScTransform function in Seurat. We used the 'scaled.data' function to identify co-expressed genes. The mean of the scaled expression was calculated for all genes in a specific cluster. The genes with the mean greater than 0 were considered as being expressed.

#### ATAC-Seq of zebrafish β-cells and hepatocytes

ATAC-Seq. was carried out using 50,000 cells collected using FACS. For FACS, single-cell suspension was generated as mentioned above. For ATAC-Seq of  $\beta$ -cells, tissue for dissociation was collected from the Tg(ins:BB1.0L; cryaa:RFP) line and β-cells sorted using the red fluorescence. Cells were processed according to the protocol outlined by Buenrostro et al. (2013). In brief, cells were homogenized in a 2 ml Dounce homogenizer with five strokes in 1× homogenization buffer on ice. The nuclei suspension was centrifuged at 500 g for 10 min at 4°C. After removing the supernatant, transposition was carried out by incubating the nuclei in the transposition reaction mix for 30 min at 37°C. Transposed DNA was purified using Qiagen MinElute Reaction Cleanup Kit. Barcoded libraries were generated from the isolated DNA by PCR amplification for 11 cycles. Libraries were sequenced on llumina HiSeq2500 in 2×75 bp paired-end mode. We conducted two biological replicates for β-cells. Processing of ATAC-seq raw data was carried out according to the pipeline from the Kundaje lab (Koh et al., 2016). MACS2 (Feng et al., 2012) was used to identify the peak regions. Only peaks present in both the biological replicates of  $\beta$ -cells were retained. The  $\beta$ -cell-specific peaks were exported as a bed file and visualized in IGV viewer (Robinson et al., 2011).

#### Transcriptional profiling and analysis of $\beta$ -, $\delta$ 1-, and hybrid cells

Collection of  $\beta$ -,  $\delta$ 1- and hybrid cells for transcriptional profiling was performed using FACS of single-cell suspension of dissected islets from Tg(ins:BB1.0L); Tg(sst1.1:EGFP-Ras). For single-cell profiling using Smart-Seq, cells were collected in 96-well plate using index sorting and processed according to the previously published pipeline (Singh et al., 2018). Analysis was performed using Seurat (Satija et al., 2015), with cells classified using their position on the FACS plot ( $\delta$ 1-cells, green<sup>+</sup> and red<sup>+</sup>; hybrid cells, green<sup>+</sup> and red<sup>+</sup>; and  $\beta$ -cells, green<sup>-</sup> and red<sup>+</sup>). For bulk-sequencing and analysis, 2000 cells of each type were collected in a 1.5 ml tube and processed according to the pipeline published by Janjuha et al. (2018). Differential gene expression analysis was analyzed using DESeq2 (Love et al., 2014).

#### Pseudotemporal analysis of single-cell RNA-seq. data

Pseudotemporal analysis was performed using Monocle 2 (Trapnell et al., 2014) following the recommended analysis protocol (http://cole-trapnell-lab.github.io/monocle-release/docs/). Briefly, raw count data was imported into a Monocle object, normalized using the estimated size factors, and filtered to remove low quality genes and cells. Gene dispersion was estimated using negative binomial distribution with fixed variance, and used to identify the highly variable genes in an unbiased manner for pseudotemporal ordering. Pseudotemporal trajectory was constructed using the 'DDRTree' algorithm, and plotted on a 1-D graph with jitter to enhance visualization. Cells in the graph were colored according to their FACS plot identity. Gene Expression dynamics in relation to pseudotime was calculated using the command 'differentialGeneTest (Monocle\_object, fullModelFormulaStr="~sm.ns(Pseudotime)")' and plotted using the heatmap function for visualization.

#### Diffusion pseudotime analysis of single-cell RNA-seq. data

For single cell trajectory inference, diffusion pseudotime was computed for 7 dpa and 14 dpa, and solely on selected cells (ins+ and sst1.1+ cells) following the standard analysis pipeline (Haghverdi et al., 2016) provided by the Scanpy package. As a single lineage of a timepoint was selected, the number of branchings was set to 1. For cell trajectory analysis, diffusion maps were developed. For 14 dpa, the sst1.1-high cells were selected as root cells to capture the transition of  $\delta1$ -cells. To visualize the marker genes trajectory on trendline plot, plot.wishbone\_marker\_trajectory function was used provided by the Wishbone package (Setty et al., 2016).

#### Tissue collection for imaging

To facilitate confocal imaging of the islets, the pancreas was dissected from fish (larval) or the gut (juvenile and adults) after fixation. Fish were killed in Tricaine prior to either direct fixation or dissection of gut, and the samples immersed in 4% paraformaldehyde+1% Triton-X for 2 days at 4°C. The pancreas was then manually dissected and washed multiple times in PBS.

## Immunofluorescence and image acquisition

Whole-mount immunofluorescence was performed on pancreas collected as described above. The collected samples were permeabilized in 1% PBT (Triton-X-100) and blocked in 4% PBTB (BSA). Primary and secondary antibody staining were performed overnight at 4°C. Primary antibodies used in this study were anti-insulin (guinea pig, Dako A0564)/insulin (rabbit, Genetex 128490) at 1:200, anti-somatostatin (rat, Genetex GTX3906) at 1:200 and anti-EGFP (chicken, Abcam ab13970) at 1:250. Secondary antibodies at 1:500 dilutions used in this study were Alexa Fluor 568 anti-guinea pig, Alexa Fluor 568 anti-rabbit, Alexa Fluor 647 anti-rabbit and Alexa Fluor 647 anti-rat. Samples were mounted in Vectashield and imaged using a Zeiss LSM 780. ImageJ was used to add scale bars and PowerPoint was used for labeling.

For immunostaining of the adult pancreatic sections obtained from controls and various time-points post  $\beta$ -cell ablation, the Tyramide Signal Amplification kit (Thermo Fisher Scientific, Alexa Fluor 555 Tyramide SuperBoost B40923) was used to amplify the signal from the insulin antibody. Primary and secondary antibody staining were performed as

recommended by the kit. Draq7 was used as the nuclear label. Sections were mounted in Vectashield and imaged using Zeiss LSM 780.

#### Live-imaging during β-cell regeneration

From 1 dpf onwards, *Tg(ins:FLAG-NTR)*; *TgBAC (sst1.1:EGFP Ras)*, *Tg (ins: BB1.0L)* triple transgenic zebrafish larvae were treated with 0.003% 1-phenyl thio-urea (PTU) (200 μM) to inhibit pigmentation. For β-cell ablation, 3 dpf larvae were incubated with 10 mM Mtz (Sigma-Aldrich, M3761) dissolved in E3 medium containing 1% DMSO and maintained for 24 h in the dark before they were rinsed and returned to fresh E3 medium. E3 medium with 1% DMSO was used as a control. Following the ablation, individual larvae were mounted in a glass-bottom micro-well dish (MakTek Corporation) in 1% low-melting agarose (LMA) containing 0.4 g/l tricaine. After the agarose was solidified, the dish were filled with E3 medium containing 0.4 g/l tricaine. Live-imaging was performed on an upright laser-scanning confocal microscope (Zeiss LSM 780). EGFP and dsRed signals were acquired simultaneously using the 488 nm and 561 nm laser lines. Laser power was maintained as low as possible to minimize photo-toxicity.

#### Quantification of bi-hormonal cell positions in the islet

Pancreatic sections from 7 and 16 mpf Tg(sst1.1: EGFP-Ras) transgenic zebrafish were co-stained with insulin and somatostatin antibodies. The number of bi-hormonal cells was quantified manually using ImageJ software. The islet periphery was determined using the 'freehand' selection tool and the islet center was computed using the 'center of mass' measurements in ImageJ (Analyze→Set Measurements→Center of Mass). The relative position of each bi-hormonal cell was determined in the following way: relative distance=distance from the islet center/distance from the islet center to the islet edge nearest to the bi-hormonal cell. The relative position of bi-hormonal cells was represented as a scatter plot with the position of the cells on the *x*-axis (islet center=0.0; islet periphery=1.0) and the age of the fish on the *y*-axis.

#### $\beta$ -Cell tracing in zebrafish larvae

From 1 dpf onwards, *Tg(ins: H2B-mEos2; cryaa: CFP)*; *Tg(ins: FLAG-NTR; cryaa: mCherry)* double transgenic zebrafish larvae were treated with 0.003% PTU (200 µM) to inhibit pigmentation. At 3 dpf, the zebrafish larvae were anaesthetized using tricaine and exposed to blue light for photoconversion. From 3 to 4 dpf, the larvae were incubated in 10 mM Mtz dissolved in E3 medium containing 1% DMSO and maintained for 24 h in the dark. Mtz was washed and the fish were returned to fresh E3 medium.

#### In vivo calcium imaging in zebrafish larvae

Live imaging and glucose injections were performed as described previously (Salem et al., 2019).

## Generation of Tg(ins:nls-mCerulean-T2A-dkk3b; cryaa:mCherry) zebrafish line

The *dkk3b* coding sequence was purchased from Zebrafish Gene Collection (ZGC) cDNA clones (MGC: 162332). The cDNA was PCR amplified to flank the cDNA with SpeI/PacI restriction sites. The PCR product was digested with SpeI/PacI restriction enzymes. The ins:nls-mCerulean; *cryaa:mCherry* construct [used for generating the *Tg(ins:nls-mCerulean; cryaa:mCherry)* line] was digested with SpeI/PacI. The two DNA fragments were ligated and transformed to generate the *ins:nls-mCerulean-T2A-dkk3b; cryaa:mCherry* plasmid. The entire construct is flanked with I-SceI sites to facilitate transgenesis. I-SceI based transgenesis was carried out, and founders selected based on mCherry (red) fluorescence in the eye and mCerulean (blue) expression in the pancreatic islet.

## Generation of Tg(Tol2-hsp70l:mCherry-T2A-dkk3b) zebrafish line

The dkk3b coding sequence was PCR amplified to flank the cDNA with NheI/BglII restriction sites. The PCR product was digested with NheI/BglII restriction enzymes and was fused to mCherry using the T2A sequence. Thereafter, mCherry-t2a-dkk3b sequence was cloned downstream of the

hsp70l promoter region. The entire construct is flanked with Tol2 sites to facilitate transgenesis. Tol2-based transgenesis was carried out, and founders selected based on ubiquitous expression of mCherry (red) fluorescence in the zebrafish larvae upon heat-shock at 37°C.

#### Generation of Tg(ins:K-GECO1; cryaa:mCherry) zebrafish line

For the construction of the *Tg(ins:K-GECO1; cryaa:mCherry)* line, we used PCR amplification with primers designed to introduce 5' EcoRI and 3' PacI restriction enzymes sites in the cDNA for K-GECO1. The established plasmid backbone containing *ins:mAG-zGeminin; cryaa:mCherry* (Ninov et al., 2013) was digested with EcoRI/PacI and K-GECO1, and the cDNA was ligated using the EcoRI/PacI sites. The construct was flanked with I-Scel sites to facilitate transgenesis. Several founders were screening and a founder with Mendelian segregation were selected for expanding the transgenic line.

#### Generation of the Tg(ins: H2B-mEos2b; cryaa: CFP) zebrafish line

For the construction of the *Tg(ins: H2B-mEos2b; cryaa: CFP)* transgenic line, the cDNA for H2B-mEos2b was PCR amplified to flank the cDNA with Mfe1/Pac1 restriction enzyme sites. A previously established plasmid backbone with *ins: mKO2-zCdt1; cryaa: CFP* (Ninov et al., 2013) was digested with EcoRI/PacI and H2B-mEos2b was ligated using the EcoRI/PacI sites (Mfe1 and EcoR1 have compatible cohesive ends). The entire construct is flanked with I-SceI sites to facilitate transgenesis. I-SceI based transgenesis was performed and founders were selected based on CFP (blue) fluorescence in the eye and green expression in the pancreatic islet.

#### **Electron microscopy**

Correlative light and electron microscopy (CLEM) of immunolabeled resin sections was performed as previously described (Fabig et al., 2012). In brief, wild-type and ablated zebrafish larvae were embedded in Lowicryl K4M using the progressive lowering of temperature method (Carlemalm et al., 1982). Ultrathin resin sections were cut on Leica UC6 ultramicrotrome, labeled with primary anti-GFP (rabbit-anti-GFP, TP401 from Torrey Pines) followed by consecutive incubations with protein A 10 nm gold, goat-anti-rabbit Alexa 488 and DAPI. Sections were analyzed on a Keyence BZ 8000 fluorescence microscope, contrasted with uranyl acetate, dried and imaged on a Jeol JEM 1400-Plus transmission electron microscope running at 80 kV.

## Statistical analysis

Statistical analysis was performed using R and GraphPad Prism. No animals were excluded from analysis. Blinding was not performed during analysis. Analysis of normal distribution was not performed. To compare the proportion of sst1.1-expressing β-cells during regeneration, an unpaired two-tailed t-test with unequal variance [t-test(x=dataframe, alternative="two.sided", paired=FALSE, var.equal=FALSE)] was used. P<0.05 was considered statistically significant. For plotting, Tukey style boxplots showing the median along with the 25th percentile (lower quartile, Q1) to 75th percentile (upper quartile, Q3) range were used. For all boxplots, whiskers extend to 1.5 times the interquartile range (IQR=Q3-Q1). In addition, individual data points are plotted on the graphs. To compare the proportions of bi-hormonal cells during time-course imaging, to evaluate the AUC during in vivo calcium imaging and the bi-hormonal cell number upon dkk3b overexpression, a Mann-Whitney unpaired t-test (non-parametric test) was used for statistical analysis. The graphs were prepared using Graph Pad prism software and P<0.05 was considered statistically significant.

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#### Competing interests

The authors declare no competing or financial interests.

#### **Author contributions**

Conceptualization: N.N., S.P.S.; Methodology: A.H., M.K., L.D.S., B.S., S.J., P.O.-C., O.K., J.B., A.K., A.P., T.K., S.R., F.R., S.E.E., J.P.J.; Formal analysis: P.C.; Investigation: N.N., S.P.S., P.C., A.H., M.K., L.D.S., B.S., S.J., P.O-C., O.K., J.B.; Resources: N.N.; Writing - original draft: N.N., S.P.S.; Supervision: N.N., J.P.J.

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#### Data availability

The processed data are available for viewing and exploration on the publicly accessible Single Cell Portal at https://singlecell.broadinstitute.org/single\_cell/study/SCP1549. Single-cell RNA sequencing data of zebrafish pancreatic cells have been deposited in GEO under accession number GSE123662 (Salem et al., 2019). Single-cell and bulk analysis datasets of pancreatic somatostatin and  $\beta$ -cells from zebrafish have been deposited in GEO under accession number GSE152697. Data for ATAC sequencing of zebrafish pancreatic  $\beta$ -cells and hepatocytes have been deposited in GEO under accession number GSE152199.

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