Single-Cell Transcriptomics of the Human Endocrine Pancreas

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Human pancreatic islets consist of multiple endocrine cell types. To facilitate the detection of rare cellular states and uncover population heterogeneity, we performed single-cell RNA sequencing (RNA-seq) on islets from multiple deceased organ donors, including children, healthy adults, and individuals with type 1 or type 2 diabetes. We developed a robust computational biology framework for cell type annotation. Using this framework, we show that α- and β-cells from children exhibit less well-defined gene signatures than those in adults. Remarkably, α- and β-cells from donors with type 2 diabetes have expression profiles with features seen in children, indicating a partial dedifferentiation process. We also examined a naturally proliferating α-cell from a healthy adult, for which pathway analysis indicated activation of the cell cycle and repression of checkpoint control pathways. Importantly, this replicating α-cell exhibited activated Sonic hedgehog signaling, a pathway not previously known to contribute to human α-cell proliferation. Our study highlights the power of single-cell RNA-seq and provides a stepping stone for future explorations of cellular heterogeneity in pancreatic endocrine cells.

The endocrine pancreas plays a critical role in the control of blood glucose homeostasis. The endocrine cells are organized into the islets of Langerhans, approximately spherical groups of 500–1,000 cells, which together constitute only 1–2% of total pancreas mass. Pancreatic endocrine cells are characterized by their most abundant hormone, namely insulin (INS; β-cells), glucagon (GCG; α-cells), somatostatin (SST; δ-cells), pancreatic polypeptide (PPY; PP cells), and ghrelin (GHRL; ε-cells). The proportion of the assorted endocrine cell types and their arrangement within the islets varies widely among different mammalian species (1). For instance, whereas rodent islets are comprised of up to 90% insulin-producing β-cells in a distinct islet core, human islets display intermingled endocrine cells, with only ~54% β-cells (2). Recently, diabetes researchers have renewed their focus on endocrine cellular heterogeneity (3). It is well accepted that not all β-cells are identical, especially in conditions of metabolic stress, such as obesity or type 2 diabetes (3–6). Moreover, it has been reported that in certain conditions of type 2 diabetes, a subset of pancreatic cells malfunction by reduction of glucose-stimulated insulin secretion or through dedifferentiation (7–9).

Individual cellular changes are diluted and therefore missed when analyzed at the level of the whole islet, or even when using sorted cell populations in bulk. Moreover, single-cell measurements can uncover unanticipated subpopulations, rare cellular states, or novel transcriptional mechanisms (10,11). Thus, methods to probe expression changes at the single-cell level are highly desirable (12–14). RNA sequencing (RNA-seq) can now be performed at the single-cell scale and, when applied in this manner, is an effective methodology for the analysis of gene expression variation among a population of apparently near-identical cells.

Here, we use single-cell RNA-seq to determine the transcriptomes of human pancreatic endocrine cells in four distinct developmental and physiological states: early childhood, normal adulthood, type 1 diabetes, and type 2 diabetes (see Fig. 1A for workflow and Table 1 for donor information). We discover that the transcriptional states of α- and

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Figure 1—A microfluidics system combined with a computational pipeline results in efficient capture of single cells from human pancreatic islets and accurate annotation of their specific cell types. 

**A:** Schematic of experimental workflow. 

**B:** Cell type annotation pipeline. Results for one β-cell are displayed as an example. For details, see RESEARCH DESIGN AND METHODS.

**C:** Abacus plot displaying the read counts per million (cpm) of key markers, including GCG (red), INS (blue), SST (orange), GHRL (green), PPY (pink), PRSS1 (brown), and KRT19 (black) in all annotated cells. Each vertical line in the abacus represents one cell. The most highly expressed marker in each cell is denoted by increased color opacity and size in relation to other markers. Within each sample, cells are ordered by GCG and thereafter INS levels. An example of expression of these markers in one cell is shown at the right of the abacus plot. In this example, the most highly expressed marker is GCG (red). All the other markers have a lower number of reads and are represented by corresponding colored dots.

**D:** Overview of cell annotation frequencies. QC, quality control; T1D, type 1 diabetes; T2D, type 2 diabetes.
β-cells are not fixed in early childhood but instead become more precisely defined as humans age. Furthermore, we find that in the diabetic state, α- and β-cells display a more immature gene signature, indicating a dedifferentiation process. Using this powerful technology, we find a high degree of gene expression variability within a given endocrine cell type and uncover Sonic hedgehog signaling as a mitogenic pathway potentially activated in replicating human α-cells.

**RESEARCH DESIGN AND METHODS**

**Human Islet Sample Acquisition and Preparation**

Human islets were acquired through the Diabetes Research Center of the University of Pennsylvania (National Institutes of Health DK-19525) and the Integrated Islet Distribution Program (IIDP; http://iidp.coh.org/). Prior to cell capture, islets were cultured in Prodo islet media (PIMS Standard) with 5% human albumin serum and a glucose concentration of 5.8 mmol/L. Islets were dissociated into single cells as described previously (15). Two integrated fluidic circuit chips, 5–10 and 10–17 μm in size (1006040 and 1006041; Fluidigm), were used for cell capture of each islet sample. The SMART-seq method was used for first-strand cDNA synthesis and PCR amplification (634833; Clontech). For two control adult donors (AAJF122 and ABAF490) (refer to Table 1 for details), ArrayControl spike-in (AM1780; Thermo Fisher Scientific) was applied during cell capture. Resulting cDNAs were pooled into a 96-well plate, and the Nextera XT DNA library preparation kit (FC-131-1096; Illumina) was used for RNA-seq library preparation according to the Fluidigm protocol (100-7168). Bulk β-cell RNA-seq data were obtained from a previous publication (16), with the addition of two samples from children.

**Sequencing**

All libraries were sequenced on the Illumina HiSeq 2500 with 100-bp single-end reads. Median read depth was 2.2 million per cell. Read alignment and gene expression quantification was performed using RNA-seq unified mapper (RUM) (17). Cells with <500,000 uniquely aligned reads were excluded from downstream analysis. A total of 82% of sequenced cells passed initial technical quality control. All sequencing data are available in the Gene Expression Omnibus (GEO) repository (accession number GSE83139; http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE83139).

**Cell Type Calling**

In light of a recent finding that the 10–17-μm C1 chip can capture a significant number of cell doublets (Fluidigm white paper), we developed a five-step pipeline to distinguish single cells from doublets and to annotate cell types. First, we crudely classified cells into different cell types based on marker genes that have well-established cell type–restricted expression. These markers include the major hormone genes (GCG, INS, SST, GHRL, and PPY), genes that encode acinar cell–specific digestive enzymes (PRSS1 and PNLP), and genes associated with ductal cells (i.e., KRT19, SPP1, and HNF1B). The expression levels of each gene were fitted by normal mixture modeling of high, low, and (or) intermediate components using the mclust R package (cran.r-project.org/web/packages/mclust) and rendered in a “violin plot.” Cells were assigned to a cell type only when they expressed specific markers in the mixture model’s high expression level component and had no conflict with other markers (Fig. 1B, violin test). Second, we derived gene expression signatures for different cell types by comparing the transcriptome of each cell type to the rest of the cells in the same endocrine or exocrine category (Fig. 1B, derivation of gene signature). Third, to further refine cell type classification, previously derived gene signatures were used to calculate the correlation of each cell to its corresponding cell type–specific signature. As expected, pure cells clustered together and showed strong correlation with only one of the signatures (Fig. 1B, correlation test). Fourth, all cells that passed these two tests were collected and their gene expression patterns rederived (Fig. 1B, refined gene
signature). Fifth, we hypothesized that any cell doublets could be explained by a linear combination of two “pure” profiles. Accordingly, the new signatures from pure cell populations were used to construct in silico mixed profiles, including all the possible combinations between different cell types. To eliminate likely doublets from further analysis, all cells were examined based on the likelihood that their profiles could be explained by an artificial in silico mixture (Fig. 1B, composition estimate). They were also inspected based on their correlation to different pure signatures (2) (Fig. 1B, correlation test). The approach outlined above is similar in concept to that used in a single-cell RNA-seq study of human islets published recently (18). However, our analysis addresses the potential issue of doublets with any combination of cell types whether they are ductal, acinar, delta, etc., and not just between α- and β-cells.

To assess background contamination levels, we sequenced 25 wells that lacked any cells by microscopic inspection after cell capture. The ratios of all transcripts versus RNA spike-ins for all sequenced wells, together with the ratio of spike-ins versus transcripts of interest (e.g., GCG) of empty wells, were modeled by mclust (cran.r-project.org/web/packages/mclust) (Supplementary Fig. 5A and B). Based on this estimate, the predicted contamination level for transcripts of interest (e.g., GCG) was calculated (Supplementary Fig. 5C) using the following equation: predicted proportion of reads from GCG contamination = [(reads from all transcripts/reads from RNA spike-ins)non-empty wells × (reads from RNA spike-ins/reads from GCG)empty wells]−1.

**Downstream Analysis**

EdgeR and DESeq2 were used with default settings for differential expression analysis (19,20). Pathway analyses were performed with Gene Set Enrichment Analysis (GSEA; http://software.broadinstitute.org/gsea/index.jsp), Ingenuity Pathway Analysis (IPA; http://www.ingenuity.com/), and the Database for Annotation, Visualization, and Integrated Discovery (DAVID; https://david.ncifcrf.gov/) (21,22). Heatmaps were generated by the “heatmap” function from the NMR R package (http://renozao.github.io/NMF), or by the “heatmap2” function from gplots R package (https://cran.r-project.org/web/packages/gplots/), with Pearson correlation as the distance function.

**RESULTS**

**Single-Cell RNA-seq Analysis of Human Islets**

We performed single-cell RNA-seq of islets from eight deceased organ donors, of whom there were three adults without diabetes, one subject with type 1 diabetes, two subjects with type 2 diabetes, and two children (Table 1). We dissociated human islets into single-cell suspensions and loaded the cell mixture onto Fluidigm C1 chips without preselection for particular cell types (Fig. 1A). The mean diameter of human pancreatic endocrine cells is 9 μm, with a range from 6 to 13 μm (data not shown). In an effort to reduce size-related sampling bias, for each donor sample, we captured cells on both the 5–10- and 10–17-μm chips. Amplified cDNAs were then indexed, pooled, and further processed to construct RNA-seq libraries (Fig. 1A). A total of 635 cells were sequenced and passed our initial sequencing quality control (Fig. 1D and Table 1).

Islets are often received after different culturing times. Since in vitro culture conditions may alter islet cell function (23–25), we assessed whether culture duration influences gene expression by comparing islet transcriptomes from a single donor with type 2 diabetes at two different time points: after 4 days in culture and after a total of 12 days in culture. Remarkably, RNA expression levels between cultured and fresh cells displayed good correlations (Spearman correlation coefficient 0.863 for α-cells and 0.857 for β-cells) (Supplementary Fig. 1). Whereas other groups have reported decreased expression of differentiation and maturity markers such as ARX, MAFA, and NEUROD1 with increased culture time (25), we observed either no change or an increase in these markers either in α- or β-cells (Supplementary Figs. 2 and 3). It is possible that the current culture conditions are improved over those used in previous studies. Furthermore, our expression profiles are based on pure α- and β-cells, whereas the previous study used laser capture microdissected bulk tissues (25). We thus conclude that differences in culture time between different donor samples have only a minimal impact on our downstream analysis.

**Cell Type Annotation**

We developed a computational pipeline for cell type annotation and for the removal of potential cell doublets (see RESEARCH DESIGN AND METHODS) (Fig. 1B). In brief, we first separated our sequenced cells into different cell types based on key marker gene expressions. Subsequently, we derived cell type–specific gene signatures for pure α-, β-, δ-, PP, ductal, and acinar cells from our samples and calculated an in silico mix of profiles for each potential combination of cell types (Fig. 1B). Next, we calculated the distance of the expression profile of each cell to all of the mixture profiles, as well as the correlation of the cell to different signatures (Fig. 1B). We successfully assigned a specific cell type to 430 cells. For a general sample overview, we plotted the marker gene expressions in each of the successfully annotated cells in an abacus plot (Fig. 1C). In total, we annotated 190 α-cells, 111 β-cells, 9 δ-cells, 18 PP-cells, 96 ductal cells, and 6 acinar cells. Due to ambiguous profiles, 205 cells were initially excluded from analysis (Fig. 1D and Table 1). A high proportion of ductal cells were obtained from the donor with type 1 diabetes, unsurprisingly since islet purity was low for this donor. Interestingly, whereas all donor samples contained some cells with undetermined annotation, most cells with an undetermined phenotype were from juvenile donors. Notably, these cells tended to have a conflicted endocrine/exocrine nature. This result could potentially be explained either by the phenomenon that islet samples from children are highly embedded in the surrounding exocrine and stromal tissue (26,27) or
by less-defined gene expression patterns in children (28,29). In contrast, most of the undetermined cells from the donors with type 2 diabetes demonstrated a mixture of endocrine cell signatures, mainly α- versus β-cells.

**Validation of Cell Type Analysis**

To evaluate the cell type classification strategy, we first plotted the aggregated single β-cell gene expression values against RNA-seq data from sorted bulk β-populations for both adult and child samples (Fig. 2A and B). We observed remarkably good agreement between the two methodologies, with a Spearman correlation of >0.80. The most abundant transcripts are INS and INS-IGF2 (Fig. 2A and B). As expected, very low-abundance genes are only detected by bulk cell expression analysis. These genes are often missed in single-cell RNA-seq experiments due to stochastic sampling during the cDNA amplification process. Interestingly, some of the discordant genes between bulk and single-cell RNA-seq are non-β-cell hormone markers, likely reflecting the limited purity of FACS-sorted human islet cells (Fig. 2A and B, labeled dots). This result demonstrates the advantage of single-cell RNA-seq for gene expression analyses of human islet cell populations, since we can retrospectively confirm cell type, thereby ensuring sample purity and eliminating gene expression signals from contaminating cells present in bulk cell fractions.

We subsequently performed hierarchical clustering and constructed a heatmap using cell type markers, important transcription factors, and signaling pathway indicators (Fig. 2C). We confirmed that well-established cell type markers were highly expressed, as expected, in their assigned cell types. For example, β-cells had high levels of PDX1, IAPP, PCSK1, MAFA, and IGF2, whereas α-cells had high expression of IRX2 and ARX. Both cell types have comparable levels of FOXA2, NKX2-2, NEUROD1, and MAFA, although cell-to-cell heterogeneity is also apparent. We further examined expression of key marker genes in all endocrine cell types plus ductal and acinar cells and found that all cells classified within a given cell type expressed high levels of their expected marker gene and low levels of all others (Fig. 2D–I and Supplementary Fig. 4). For example, β-cells expressed high levels of INS and low levels of acinar and ductal markers, whereas duct cells showed high transcript levels for KRT19 but only marginal signals for hormone genes (Fig. 2D–I and Supplementary Fig. 4). Furthermore, our data show a high degree of similarity to recently published single-cell transcriptomic data of 60 pancreatic islet cells from control adult donors (18). For example, ARX is expressed highly in both α- and PP cells, whereas ETV1 is highly enriched in PP and δ-cells (Supplementary Fig. 4). Together, these observations demonstrate the high accuracy of our cell type annotation methodology.

**Assessment of Background Contamination Levels**

Some of the assigned cells demonstrate mRNA expression of markers not normally associated with that cell type. For example, low levels of GCG expression were also noted in non-α-cells (Fig. 1C, faint, small red dots). Since we rigorously excluded cells that could represent doublets, we next examined whether the secondary markers could be a result of RNA contamination originating from lysed cells prior to capture. To this end, we sequenced cDNA from 25 empty wells and modeled the relationship between the number of all mapped transcripts, reads from spike-in RNA, and reads from GCG transcripts to predict the level of GCG mRNA expression arising from contamination (see *Research Design and Methods* and Supplementary Fig. 5). Based on this analysis, we concluded that the low levels of GCG found across all cell types most likely resulted from contamination.

**Gene Expression Profiles of α- and β-Cells From Donors With Type 2 Diabetes Show Features of Juvenile Gene Activation**

Next, we derived α-cell and β-cell gene signatures from adult donors without diabetes. By design, when these genes (indicated as black bars in Fig. 3A and B) are projected back on the rank-ordered differentially expressed genes between these two cell types, α-cell signature genes cluster on the α-cell side (Fig. 3A), and β-cell signature genes are expressed only in β-cells (Fig. 3B). When we then analyzed the expression pattern of these adult signature genes in α- and β-cells from young children, many of them were no longer expressed in the expected cell type–specific fashion (Fig. 3C and D). Multiple α-cell signature genes were in fact preferentially expressed in juvenile β-cells (Fig. 3C). This inappropriate gene activation profile is even more pronounced for α-cells of young children; a very large fraction of adult β-cell genes are expressed at high levels in α-cells (Fig. 3D). Consequently, the gene set enrichment scores for the adult endocrine cell gene signatures are much lower in the young α- and β-cells, indicative of an only partially complete differentiation program in juvenile islets. The lists of genes that are misexpressed in children are presented in Supplementary Table 1. For lists of genes used to define α- and β-cell signatures see Supplementary Table 2.

Subsequently, we analyzed the enrichment of the non-diabetic adult islet cell gene signatures in α- and β-cells from individuals with type 2 diabetes. Gene sets from donors with type 2 diabetes had intermediate enrichment scores between those from adult control subjects and children (Fig. 3E and F). Similarly, several of the α- or β-cell signature genes were expressed at high levels in the inappropriate endocrine cell type (Fig. 3E and F). This feature is reminiscent of the pattern seen when comparing gene sets between child donors and adult control donors (Fig. 3C and D). This resemblance of endocrine cells between type 2 diabetic samples and child samples was confirmed when we derived α- and β-cell gene signatures from our juvenile organ donors and compared them to the gene expression patterns present in normal and type 2 diabetic adult endocrine cells (Fig. 4). Again, gene expression profiles of both α- and β-cells from donors with type 2 diabetes exhibited features of gene activation seen in children (Fig. 4A).
Figure 2—Single-cell RNA-seq of human islets identifies pancreatic cell types and gene signatures. Single-cell expression data from β-cells were aggregated and compared with expression profiles from sorted bulk β-cells in adult (A) and child (B) samples. Outlier genes indicative of contaminating cells in the bulk samples are labeled. FPKM, fragments per kilobase of transcript per million. C: Heatmap showing hierarchical clusters of α- and β-cells from different donor types with selected genes. D–I: Violin plots confirm that annotated cell types have mutually exclusive marker gene expressions. Each dot inside the violin represents one cell. T2D, type 2 diabetes.
and B). In β-cells, these include cell cycle and regulatory genes such as CDKN2B, BARD1, and JUNB (Supplementary Table 1); at least one gene involved in insulin secretion, PRKD1, is also upregulated in type 2 diabetic β-cells (30). This finding suggests that endocrine cells in patients with type 2 diabetes are not able to maintain a fully differentiated gene expression profile, which is in line with recent publications reporting partial dedifferentiation of β-cells in diabetic states (8,31). The complete list of misexpressed genes can be found in Supplementary Table 1.

**Human Proliferating α-Cells Activate the Sonic Hedgehog Pathway**

One of the α-cells analyzed from an adult donor without diabetes was clearly proliferating, as indicated by a very high level of Ki67 mRNA expression. We compared the expression profile of this proliferating α-cell with quiescent

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**Figure 3**—Child α- and β-cells exhibit a different transcriptome profile than adult α- and β-cells. Adult gene signatures for gene set enrichment analysis were derived from control adult endocrine cells with a ≥10-fold difference between α- and β-cell gene expression levels and a false discovery rate < 5%. Differentially expressed genes from control adult α- and β-cells ordered by fold changes with genes more highly expressed in α-cells on the left of the x-axis in A and genes more highly expressed in β-cells on the left of the x-axis in B. C–F: Similar display as in A and B, except that the same nondiabetic adult gene signature was used to analyze α- and β-cells from child donors (C and D) and donors with type 2 diabetes (E and F). T2D, type 2 diabetes.
α-cells from the same donor and performed pathway analysis. We confirmed that this cell had, in fact, activated cell cycle pathways and inhibited cell cycle checkpoint control (Fig. 5A). Remarkably, the Sonic hedgehog signaling pathway was activated in the replicating α-cell (Fig. 5A). DYRK1A and GSK3B were both repressed in this proliferating cell, consistent with recent reports that inhibition of both molecules induces proliferation of human endocrine cells (32,33) (Fig. 5B). Interestingly, both of these proteins regulate GLI transcription factors, major targets of Sonic hedgehog pathway signaling (Fig. 5B). The ability to analyze differential expression between a proliferating α-cell and quiescent α-cells highlights the power of single-cell RNA-seq to explore rare cell populations.

**Deeper Analysis of Cells With Conflicted Expression Profiles**

Our original exclusion of cells with a mixed signature was designed to eliminate any cells captured as doublets. This rigorous exclusion could have the detrimental effect of omitting interesting cells, such as transdifferentiating cells and even possibly rare progenitor cells. For example, recent publications have demonstrated the plasticity of pancreatic endocrine cells in transition from one cell fate to another under experimental or pathophysiological conditions (8,34–36). This plasticity corresponds to the highly similar chromatin and epigenetic state of human α- and β-cells (16,36). We therefore re-evaluated cells that were excluded from our analysis based on a conflicted α/β-expression profile by comparing their gene expression pattern to that of annotated α- and β-cells, using hierarchical clustering of both genes and cells (Fig. 6A). As seen in Fig. 6A, many conflicted cells with a mixed α-/β-signature clustered together on the heatmap in between α- and β-cells and displayed both α- and β-cell signatures at roughly equal strength (cells in a black box); they thus represent true doublets. However, several of the conflicted cells did not cluster with these doublets but rather with either α- or β-cells (see red arrows on top of the heatmap) and thus likely represent rare endocrine cells that were excluded on the very stringent criteria we described above.

We also examined α-, β-, and conflicted α-/β-cells using principle component analysis based on the same differentially expressed genes (Supplementary Fig. 6A). There are distinct three clusters of cells, corresponding to pure α- and β-cells and the group of cells showing mixed α-/β-signature, based on the first two principle components, PC1 and PC2. Strikingly, both PC1 and PC2 show high correlation with the number of transcripts sequenced (Supplementary Fig. 6A). The observation that many of the suspected admixed cells have relatively high numbers of transcripts further corroborates the notion that they are likely to be doublets (Supplementary Fig. 6A). However, we again observed cells with conflicted gene expression signatures that do not cluster with the three major groups of cells, representing potential rare cell types (Supplementary Fig. 6A).

Similarly, we performed this analysis for conflicted β-/duct cells (Fig. 6B). This analysis demonstrates that in addition to the conflicted cells that are clearly the results of two cells in one well, there are other rare cells that do not display the standard α-, β-, or duct cell signature that are not doublets. Future research with larger sample sizes will likely reveal interesting properties about rare cell types.

**DISCUSSION**

Pancreatic transcriptome studies of bulk-sorted endocrine cell fractions have revealed important insights into cell type-specific gene expression signatures (36,37). In our study, we obtained RNA-seq data from well-annotated single cells from multiple organ donors covering children, healthy adults, and individuals with type 1 and type 2 diabetes. We developed a robust pipeline to ensure that
each cell is correctly assigned to a specific cell type and that artifacts and cell doublets were removed. We also demonstrate several clear advantages in analyzing data from single cells versus the entirety of a population, even when cells are FACS sorted prior to RNA extraction.

A particularly striking result from our study is the discovery that the transcriptomes of both α- and β-cells from donors with type 2 diabetes exhibit features of expression profiles of their juvenile counterparts, indicating partial regression to an immature state, as exhibited by upregulation of several cell cycle genes (Figs. 3 and 4 and Supplementary Table 1). This was made possible by the derivation of α- and β-cell signature gene lists derived from 100% pure cells identified by single-cell technology, which is not possible through bulk cell sequencing analysis. The mechanisms driving this dedifferentiation will need to be explored further, although the transcription factor FOXO1 clearly plays a role in this process (8).

A second major advantage of single-cell technology is the ability to detect rare cell populations. We were able to discover a single proliferating α-cell, thus showcasing the power of single-cell RNA-seq methodology (Fig. 5). We demonstrate the presence of the expected activation of cell cycle genes, as well as downregulation of both DYRK1A and GSK3B in this replicating cell, both of which contribute to endocrine cell replication (32,33) and

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**Figure 5**—Sonic hedgehog signaling may contribute to endocrine cell replication. **A:** A replicating α-cell found by single-cell RNA-seq analysis showed activation of the cell cycle and repression of checkpoint control genes and, unexpectedly, activation of the Sonic hedgehog pathway. **B:** Differentially expressed genes in the Sonic hedgehog pathway are depicted with a circle with color-filled interior (red, upregulation; green, downregulation in the replication α-cell, compared with quiescent α-cells from the same donor).
therefore implicate a role for Sonic hedgehog signaling in this process (Fig. 5).

Furthermore, we performed a detailed analysis of the cells that showed a conflicted signature based on their individual expression profiles. While many clustered together, some of these cells with a somewhat "mixed" gene expression signature clustered within the bona fide α- and β-cells, or duct cells and thus possibly represent transitional states. Further research is required to uncover the biological significance of these cells.

In conclusion, by applying single-cell RNA-seq technology to profile pancreatic islets from multiple human organ donors, comparing both pediatric and adult organ donors with and without diabetes, we uncover several previously unknown features of the human endocrine pancreas. First, gene expression even of the major hormone genes is quite variable among human α- and β-cells. Second, α- and β-cells from type 2 diabetes exhibit transcriptome features of pediatric endocrine cells, indicating a reversal to a more immature state. In light of our recent findings that aged mouse islets have enhanced insulin secretion compared with young islets, the similarity of type 2 diabetic and young α- and β-cells may indicate that type 2 diabetes could in part result from a failure of compensatory mechanisms that normally occur with age (29). To what degree these abnormalities contribute to the etiology of diabetes will have to be established in the future; nevertheless, the data presented here illustrate

Figure 6—Heatmaps help distinguish doublets from rare variant cells. A: Annotated α- and β-cells, together with suspected α- and β-doublets, are clustered using genes that are >6.5-fold differentially expressed between α- and β-cells. Each column represents one cell and each row represents one gene. Cells from different sample types are color coded and indicated by the color bar on top of the heatmap. Cells with an ambiguous signature are depicted with a red arrow. Many ambiguous cells cluster together between α- and β-cells and display clear evidence of a combined signature (black box). Other ambiguous cells, however, exhibit expression profiles more closely related to either the α- or β-cell gene expression pattern and thus could represent rare transitional cells. B: Similar organization as in A, except that annotated β-cells and duct cells, together with suspected β-cell and duct cell doublets, are clustered using genes that are >3.2-fold differentially expressed between β-cells and duct cells. T1D, type 1 diabetes; T2D, type 2 diabetes.
the power of a well-controlled single-cell transcriptomic analysis.

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