Human islets contain a subpopulation of glucagon-like peptide-1 secreting α cells that is increased in type 2 diabetes

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ABSTRACT

Objectives: Our study shows that glucagon-like peptide-1 (GLP-1) is secreted within human islets and may play an unexpectedly important paracrine role in islet physiology and pathophysiology. It is known that α cells within rodent and human pancreatic islets are capable of secreting GLP-1, but little is known about the functional role that islet-derived GLP-1 plays in human islets.

Methods: We used flow cytometry, immunohistochemistry, perfusions, and calcium imaging techniques to analyse GLP-1 expression and function in islets isolated from cadaveric human donors with or without type 2 diabetes. We also used immunohistochemistry to analyse GLP-1 expression within islets from pancreatic biopsies obtained from living donors.

Results: We have demonstrated that human islets secrete ~50-fold more GLP-1 than murine islets and that ~40% of the total human α cells contain GLP-1. Our results also confirm that dipeptidyl peptidase-4 (DPP4) is expressed in α cells. Sitagliptin increased GLP-1 secretion from cultured human islets but did not enhance glucose-stimulated insulin secretion (GSIS) in islets from non-diabetic (ND) or type 2 diabetic (T2D) donors, suggesting that β cell GLP-1 receptors (GLP-1R) may already be maximally activated. Therefore, we tested the effects of exendin-9, a GLP-1R antagonist. Exendin-9 was shown to reduce GSIS by 39% and 61% in ND islets and T2D islets, respectively. We also observed significantly more GLP-1+ α cells in T2D islets compared with ND islets obtained from cadaveric donors. Furthermore, GLP-1+ α cells were also identified in pancreatic islet sections obtained from living donors undergoing surgery.

Conclusions: In summary, we demonstrated that human islets secrete robust amounts of GLP-1 from an α cell subpopulation and that GLP-1R signalling may support GSIS to a greater extent in T2D islets.

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Keywords Human islets; Glucagon-like peptide-1; GLP-1; Glucagon; Alpha-cell; Beta-cell

1. INTRODUCTION

It is known that α cells within rodent and human pancreatic islets are capable of secreting glucagon-like peptide-1 (GLP-1) [1–3] under certain conditions and the enzyme responsible for GLP-1 cleavage and subsequent inactivation, dipeptidyl peptidase-4 (DPP4), is expressed in α cells within human islets [4,5]. Despite the evidence for the production and inactivation of GLP-1 in islets, the functional role that islet-derived GLP-1 might play in human islets remains unclear.

GLP-1 therapies such as DPP4 inhibitors and GLP-1R peptide agonists have been developed and are now in wide clinical use using the classical incretin effect as a physiological framework [6]. The incretin effect describes stimulated GLP-1 release from the entero-endocrine L cells of the gut to act on pancreatic β cells, thus potentiating insulin secretion in response to nutrient ingestion [7]. Early reports have shown proglucagon cleavage to be tissue-specific with GLP-1 being produced in the L cells and negligible amounts found in islets. In contrast, glucagon secretion is limited to pancreatic islets with little if any being produced from the L cells [8–10]. However, there have also been reports of GLP-1 secretion from islets [11,12] and it has been proposed that GLP-1 may also act as a paracrine factor in islets [2,13,14], whereby α-cell GLP-1 secretion, dependent on prohormone convertase 1/3 (PC1/3) proglucagon cleavage, acts on adjacent β cells to augment function and insulin secretion. Despite the controversy regarding a functional role for intra-islet GLP-1, this paracrine concept has received advanced support from genetic mouse studies showing intra-islet GLP-1 is required for whole-body glucose homeostasis [15,16]. Although the physiological relevance of intra-islet GLP-1 in these genetic models has been demonstrated, it is currently unknown whether intra-islet GLP-1 is relevant for human physiology and pathophysiology such as T2D, and little information is available as to whether this proposed GLP-1 paracrine axis functionally exists in human islets.

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Human islet architecture argues for perhaps a greater functional paracrine role for intra-islet GLP-1 than in mouse islets because the α cells within human islets are greater in number and interspersed throughout the islet compared with the mouse islet where α cells form a mantle around the periphery [17]. Consequently, human α cells are organized to facilitate paracrine interactions with other islet cells [10]. Moreover, α cell glucagon secretion has been reported to determine the glycemic set-point in mice and humans, suggesting an important integration of α and β cell communication in glucose homeostasis [19]. As a result of the paucity of functional information on intra-islet GLP-1 from human tissue, we therefore studied isolated human islets from ND and T2D cadaveric donors as well as pancreatic biopsy sections obtained from living donors during surgery.

2. METHODS

2.1. Mouse pancreatic islets
All animal studies followed the guidelines issued by the University of Alberta Animal Care and Use Committee (protocol # A000286). Pancreatic islets from male C57BL/6 mice (ages 12–24 weeks) were isolated by collagenase Type V (Sigma–Aldrich) digestion of the pancreas and purified by Histopaque density gradient. Mouse islets were cultured in RPMI 1640 media (11.1 mM glucose) supplemented with 10% FBS and 1% penicillin/streptomycin.

2.2. Human islets and pancreatic sections
De-identified human primary islets isolated from deceased donors were obtained from the Alberta Diabetes Institute IsletCore and the Clinical Islet Laboratory, University of Alberta, in accordance with institutional human ethics guidelines. Details on the islet and donor parameters are provided in the supplementary material. If the islet preparation was <90% pure, islets were handpicked to obtain >90% purity. Islets were cultured in DMEM (5.5 mM glucose) supplemented with 10% FBS and 1% penicillin/streptomycin for static incubations and perifusions. For flow cytometry and live imaging, islets were cultured in CMRL medium (5.5 mM glucose) supplemented with 0.5% BSA, 1% Insulin-Transferrin-Selenium (ITS), 1% Glutamax, and 1% penicillin/streptomycin. Sections of isolated human islets and pancreatic sections were formalin fixed (Z-fix), paraffin embedded, and sectioned at 5-μm thick. Diagnosis of type 2 diabetes was determined from medical records and an HbA1c > 6.5%. Whole pancreatic sections from cadaveric donors were obtained from the Alberta Diabetes Institute IsletCore. Human pancreatic biopsies were generously provided by Dr. Marko Barovic and Dr. Michele Solimena from the Paul Langerhans Institute, Dresden, Germany for islet sections from living donors. Live-donor biopsies were collected from type-2 diabetic and non-diabetic patients undergoing treatment for either chronic pancreatitis or malignant pancreatic tumors. Biopsies were acquired from the bodies and tails of the intact pancreases during laparoscopic surgery, or from the bodies of excised whole pancreases after a pancreatectomy. Immediately after surgery, the biopsies were fixed in 4% PFA at room temperature for 2–3 h. After fixation, the tissue samples were thoroughly washed and stored in PBS with added 0.02% sodium azide at 4 °C until paraffin embedding. Samples were paraffin-embedded and sectioned at the Paul Langerhans Institute before shipment to the University of Alberta.

2.3. Antibodies for immunofluorescent microscopy
Anti-glucagon Rab [EP370] (ab92517) and Anti-GLP-1 (amidated) Mab [BG9] (ab26278) were purchased from Abcam. Anti-insulin guinea pig polyclonal DAKO (A056401-2) was purchased from Agilent Technologies. Alexa Fluor goat anti-rabbit 488, Alexa Fluor goat anti-mouse 568, and Alexa Fluor goat anti-guinea pig 647 were used as secondary antibodies and purchased from Thermofisher Scientific.

2.4. Antibodies for flow cytometry
For islet culture experiments, sitagliptin (50 ng/ml) or IL-1β (10 ng/ml) were added to KRBH buffer containing 2.5 mM glucose, 25 islets were perfused at 250 μL/min with 2.5 mM glucose buffer, 25 islets were perfused at 250 μL/min with 2.5 mM glucose buffer for 20 min. Next, islets were perfused in 11.1 mM glucose buffer, 25 islets were perfused at 250 μL/min with 2.5 mM glucose buffer for 20 min. Next, islets were perfused in 11.1 mM glucose buffer, 25 islets were perfused at 250 μL/min with 2.5 mM glucose buffer for 20 min. Next, islets were perfused in 11.1 mM glucose buffer, 25 islets were perfused at 250 μL/min with 2.5 mM glucose buffer for 20 min.
Figure 1: Human islets contain a GLP-1 secreting α cell subpopulation. A. Human islets secrete ~50-fold more active GLP-1 than mouse islets. B. Representative images of human islets stained for glucagon, GLP-1, and double-positive staining (merge, arrows denote examples of double-positive cells) indicating a subpopulation of α cells. Scale bar: 20 μm. C. Quantification of glucagon+ cells shown as % total islet cells. N = 6 donors. D. Quantification of amidated GLP-1+ cells shown as % total glucagon+ cells. N = 6 donors. E. Representative merged image of amidated GLP-1/glucagon staining from a section of whole human pancreas (arrows denote examples of double-positive cells). Scale bar: 50 μm. F. Representative flow cytometry contour plots of dispersed islet cells. Left, insulin and glucagon expression were analysed in total dispersed islets, allowing the identification of insulin-/glucagon-, insulin+, and glucagon+ cell populations. Right, the insulin-/glucagon-, insulin+, and glucagon+ cell populations were further analysed for forward scatter (FSC) and expression of amidated GLP-1. Amidated GLP-1+ cells were gated within each cell population. G. Percentages of amidated GLP-1+ cells in the insulin-/glucagon-, insulin+, and glucagon+ cell populations. N = 6 donors. H. Representative flow cytometry histograms showing PC1/3 expression levels within GLP-1-/glucagon-, GLP-1+/glucagon+, and insulin+ cells. I. Relative expression of PC1/3 expression in GLP-1-/glucagon+ cells, GLP-1+/glucagon+ cells, and insulin+ cells. Median fluorescent intensities (MFIs) of PC1/3 staining were calculated within each cell fraction, normalized relative to the insulin+ cell fraction, and compiled as percent expression to insulin+ cells. J. Representative flow cytometry histograms showing DPP4 expression in insulin-/glucagon-, insulin+, and glucagon+ cells. K. Relative expression of DPP4 expression in insulin-/glucagon-, insulin+, and glucagon+ cells. MFIs of DPP4 staining were calculated within each cell population, were normalized relative to the insulin-/glucagon-cell fraction and compiled as fold expression relative to insulin-/glucagon-cells. N = 3 donors. L. Islet section antibody validation immunoblot showing that the antibody used in this study to detect amidated forms of GLP-1 does not detect glucagon. Statistical significance for the data was determined using a paired Student’s t-test. *, P < 0.05; ***, P < 0.001. Error bars indicate SEM.
glucose buffer for 50 min. Perfusate was collected at timed intervals and stored at –20 °C for insulin analysis.

2.8. Calcium imaging

Human pancreatic islets were incubated with 2 μM Fluo-4 AM for 1 h in a humidified incubator at 37 °C and 5% CO2. They were placed in the recording chamber of an Olympus IXG3 inverted fluorescence microscope equipped with an UPLANApo 10X objective. We used a multichannel gravity-fed Warner Instruments TC-324B temperature-controlled perfusion system, maintained at 37 °C. Excitation was performed with a X-Cite 120LEDBoost (Excellitas) light source and image acquisition was performed with an Andor iXon ultra camera using Olympus cellSens software. The fluorescence of whole human pancreatic islets was measured at a sampling frequency of 0.33 Hz. The islets were perfused with a solution containing (mM) 120 NaCl, 4.8 KCl, 1.2 MgCl2, 2.5 CaCl2 and 10 HEPES (pH 7.4) supplemented with the indicated amount of glucose and Exendin-9. Depolarization of the islets with 30 mM KCl was used as a positive control at the end of the measurement. Ca2+ events and signal dynamics of active subregions within the islets, before or during the application of Exendin-9, were analysed.

2.9. Hormone secretion assays

Active GLP-1 levels from islet culture media samples (human and mouse) were quantified using the electrochemiluminescent assay active GLP-1 (v2) kit (K150JWC-1), MesoScale Discovery (MSD). The MSD active GLP-1 (7–36) antibody was validated in-house against high purity synthetic peptides and was found to be highly selective for the GLP-1 (7–36) isoform when compared with GLP-1 (9–36), GLP-1 (1–36), and glucagon (Figure 2F). Total GLP levels in human islet

![Figure 2](https://example.com/figure2.png)

**Figure 2:** Sitagliptin increases active GLP-1 levels in cultured islets, but does not increase GSIS from ND and T2D islets. **A.** Active GLP-1 secretion from human islets in culture showing the inhibitory effects of sitagliptin on DPP4 over 24 h. N = 4 donors. **B.** Active GLP-1 secretion from human islets treated with sitagliptin in serum-free culture (5.5 mM glucose) for 48 h to exclude the effect of serum DPP4. N = 3 donors. **C.** Active GLP-1 secretion increases with sitagliptin treatment in control media or IL-1 beta (50 ng/ml) co-incubation. N = 3 donors. **D.** Sitagliptin is unable to increase insulin secretion with glucose-stimulated perifusions of ND islets. 25 islets per lane, N = 6 donors. **E.** Sitagliptin is unable to increase insulin secretion with glucose-stimulated perifusions of T2D islets. 25 islets per lane, N = 3 donors. For all experiments the sitagliptin concentration used (200 nM) was in the therapeutic range. **F.** GLP-1 antibody validation plot illustrating that the GLP-1 assay used in this study is highly selective for active GLP-1 compared with the inactive GLP-1 peptides (9–36) and (1–36) and glucagon. Statistical significance for the data was determined using a paired Student’s t-test or two-way ANOVA. *P < 0.05, **P < 0.01, ***P < 0.001. Error bars indicate SEM.
culture media samples were quantified with the electrochemiluminescent assay MSD total GLP-1 (v2) kit (K150JVC-1). Insulin levels in supernatants from static incubations and perfusate samples were quantified with Stellux Chemiluminescent Human Insulin ELISA, ALPCO.

2.10. Immunofluorescence microscopy
Paraffin-embedded human islet sections were rehydrated and subjected to antigen retrieval by microwaving the slides for 15 min in a citrate buffer (pH 6.0). After cooling, sections were blocked with 20% normal goat serum for 1 h, incubated with primary antibodies for 1 h, and then incubated with secondary antibodies and DAPI (1:1000) for 1 h. ProLong Gold antifade mountant was applied and the slide sections were mounted on coverslips. Primary antibodies were as follows: 1:1000, anti-glucagon Rab [EP370] (ab92517), anti-insulin guinea pig polyclonal DAKO (A056401-2); 1:2000 anti-GLP-1 amidated Mab (B8G9) (ab26278)). This amidated GLP-1 antibody was validated in house and detected the C-terminal amidated 1–36, 7–36, and 9–36 GLP-1 isoforms but not glucagon (Figure 1L). Secondary antibodies were incubated at 1:200 dilution. For epifluorescence microscopy, islet sections were imaged with an Olympus IX83 inverted fluorescence microscope and a UPlanSapo 20X objective. Excitation was performed with an X-Cite 120LEDBoost (Excellitas) light source with the appropriate filter set for DAPI, AF488, AF568, and AF647. Images were captured with a Hamamatsu Orca Flash4.0 camera operated with Olympus cellSens software. Image analysis was performed using Fiji and a customized macro. DAPI staining was used to identify ROIs (cells). Where appropriate, thresholds were set to identify positive staining in the green (AF488), red (AF568), or far-red (AF647) channels.

3. RESULTS
3.1. Human islets contain a GLP-1 secreting α-cell subpopulation
As human islets possess ~40% α cells compared with ~20% α cells in mouse islets [17], it is reasonable to expect that human islets would secrete ~2-fold more active GLP-1 when compared with mouse islets. However, we observed a ~50-fold increase in active GLP-1 in human islet cultures compared with mouse islets suggesting that GLP-1 secretion from α cells may be more important to human islet function (Figure 1A). Because the number of α cells is greater and GLP-1 secretion is higher in human islets, we hypothesized that a significant α-cell subpopulation would express the post-processed amidated form of GLP-1. Human islet sections were stained with antibodies for glucagon and amidated GLP-1. The amidated GLP-1 antibody was validated for selectivity using an in-house immunoblot assay (Figure 1L) that shows that the antibody can detect amidated forms of GLP-1, but not glucagon. We determined that 40% of the total cells in human islets are α cells and that ~40% of the α cells are a subpopulation that is positive for GLP-1 (Figure 1B–D). Importantly, GLP-1 expression was also found in a subpopulation of α cells in pancreatic biopsies, indicating that the presence of intra-islet GLP-1 is not an artifact of islet isolation and culture (Figure 1E). To study this subpopulation further, flow cytometry was used to segregate three cell populations within human islets: insulin+/glucagon−, insulin+, and glucagon+ (Figure 1F). Our results show that GLP-1 expression is found only in a subpopulation of glucagon+ α cells that constitutes ~50% of the total α cell number (Figure 1F,G), a value that is consistent with the value of 40% observed in human islet sections (Figure 1D). PC1/3 is required for the proteolytic cleavage of GLP-1 from proglucagon and we observed a significant increase in PC1/3 expression in glucagon+/GLP-1+ α cells compared with glucagon+/GLP-1− cells (Figure 1H,I). We further characterized the α cell population using DPP4 expression and confirmed that DPP4 expression is enriched in α cells [4,5] when compared with β cells (Figure 1J,K). However, we did not observe any differences in DPP4 expression between GLP-1+ and GLP-1− α cells (data not shown).

3.2. Sitagliptin increases active GLP-1 levels in culture, but does not enhance GSIS in ND and T2D islets
Because human α cells express DPP4, we tested the ability of sitagliptin (a DPP4 inhibitor) to increase active GLP-1 levels in human islet cultures. We observed a significant increase in active GLP-1 levels with sitagliptin treatment in long-term cultures (24–48 h, Figure 2A,B,C). We also tested the ability of sitagliptin to increase active GLP-1 in a serum-free culture and confirmed that intra-islet DPP4 is being inhibited, rather than any DPP4 that may be in the culture media (Figure 2B). As inflammatory cytokines may increase GLP-1 secretion, we tested the effects of IL-1β, however the sitagliptin-mediated increase in active GLP-1 was not affected by that cytokine challenge (Figure 2C). As sitagliptin increased active GLP-1, we hypothesized that sitagliptin would augment GSIS. However, under perfusion conditions, sitagliptin-treated ND islets did not show any increase in GSIS (Figure 2D). Because DPP4 inhibitors were successfully used to increase circulating active GLP-1 levels and restore insulin secretion in T2D patients [20], we tested whether sitagliptin would enhance GSIS in T2D islets. Again, no increase in insulin secretion was observed in perfused GSIS experiments, indicating that DPP4 inhibition in T2D islets does not directly contribute to increased GSIS (Figure 2E). To validate the selectivity of the active GLP-1 assay used in these studies, we tested the active GLP-1 (7–36) antibody against several other proglucagon cleavage peptides. We observed that the antibody was highly selective for the active GLP-1 (7–36) peptide against several other proglucagon cleavage peptides. We observed that the antibody was highly selective for the active GLP-1 (7–36) peptide, whereas no signal was detected with either the (1–36) or (9–36) GLP-1 peptides or glucagon (Figure 2F).

3.3. Islet GLP-1Rs contribute to insulin secretion in ND and T2D islets
Because sitagliptin treatment did not enhance GSIS in ND and T2D islets, one plausible explanation is that the GLP-1Rs may be maximally stimulated by the robust amount of intra-islet proglucagon peptides that are present, such as GLP-1 (Figure 1A). To explore this concept, we used the GLP-1R antagonist, exendin-9, under the GSIS perfusion conditions and observed a 25% decrease in insulin secretion that occurs primarily at high glucose in ND islets (Figure 3A), suggesting that GLP-1R signalling contributes to insulin secretion in ND islets. In T2D islets, we observed an even larger decrease in insulin secretion of 62% that occurred at both low- and high-glucose conditions (Figure 3B), suggesting that insulin secretion is strongly supported by GLP-1R signalling in T2D. Indeed, when the change in insulin secretion over time is analysed for ND and T2D islets, the percent change from control is 61% greater in T2D islets than in ND islets (Figure 3C). As fluctuations in intracellular calcium correlate with β-cell insulin secretion, we investigated whether treating human islets with exendin-9 would alter calcium dynamics. ND islets were loaded with the calcium sensitive indicator Fluo-4 and stimulated with glucose (10 mM). Exendin-9 caused a reduction in calcium dynamics in whole islets as well as dynamic sub-regions of islets (Figure 3D,E). Calcium events and signal dynamics of sub-regions were significantly increased.
decreased in the presence of Exendin-9 when compared with the high (10 mM) glucose control (Figure 3F, G).

3.4. The GLP-1 expressing α cell subpopulation is increased in T2D islets

Because GLP-1 expression in pancreatic α cells has been associated with metabolic stress and β cell injury [21, 22], we hypothesized that the GLP-1+ α cell subpopulation may be increased in T2D. To test this concept, we obtained islet sections from T2D and ND cadaveric donors and stained for glucagon and amidated GLP-1. Double positive staining for glucagon and GLP-1 was observed in both ND and T2D islets (Figure 4A, B). Image analysis revealed that the GLP-1+ α cell subpopulation was significantly increased (Figure 4C). We then immuno-stained islet sections from the same donors for glucagon and insulin to identify the α and β cell populations, respectively (Figure 4E, F). Image analysis of T2D islets showed no change in α cell density, but a significant decrease in β cell density (Figure 4G, H) resulting in a
significant decrease in the insulin+/glucagon + cell ratio, confirming that the islets are phenotypically diabetic (Figure 4I).

3.5. The GLP-1 expressing α-cell subpopulation is present in islets obtained from living donors

Although intra-islet GLP-1 has previously been detected in human and rodent islets, it has been suggested that this is because of stress-induced alterations in proglucagon peptide processing resulting from ischemia, isolation, and culturing. Therefore, it is important to determine whether intra-islet GLP-1 is present in human islets that have not undergone these stressors. Accordingly, we analyzed pancreatic sections obtained from living donors undergoing surgery for pancreatitis or malignancy because the presence of intra-islet GLP-1 in those sections would provide further evidence for a physiological role in humans. Analysis of pancreatic sections from eleven living donors revealed that the islets had a similar percentage of insulin+ β cells and insulin+/glucagon + cell (β/α) ratio (Figure 5A–C) to islets obtained from cadaveric donors (Figure 4H). Furthermore, immunostaining for glucagon and amidated GLP-1 (Figure 5D–F) revealed that ~70% of the glucagon + cells were also positive for GLP-1 (Figure 5G). Four of the eleven living donors had type 2 diabetes, so we compared insulin+, glucagon+, and GLP-1+ cell populations between non-diabetic donors and those with type 2 diabetes. As was previously seen in islet sections from cadaveric donors (Figure 4H), we also observed a significant reduction in insulin + cells in type 2 diabetic islets from living donors (Figure 5H). However, in contrast to our
Figure 5: A subpopulation of GLP-1+ / Glucagon + (x) cells can be detected in islets within pancreatic sections obtained from living donors undergoing pancreatic surgery. A. Representative images of glucagon + and insulin + staining from a single human islet. B,C. Pancreatic section grouped data showing (B) the number of insulin + cells as a % of the total number of cells (DAPI staining in blue) and (C) the ratio of insulin + to glucagon + cells. D. Representative images from a single human islet showing glucagon +, GLP-1 + staining and their co-localization (merge, arrows denote examples of double positive cells). E-G. Pancreatic section grouped data showing the number of: (E) Glucagon + cells and (F) GLP-1 + / glucagon + double positive cells as a % of the total number of cells (DAPI staining in blue), (G) GLP-1 + cells as a % of the total number of glucagon + cells. H. Grouped data from pancreatic sections obtained from non-diabetic (ND) and type 2 diabetic (T2D) living donors. Islets were stained for insulin, glucagon, and GLP-1. I. Grouped data expressed as a % of the number of GLP-1 + cells of the total glucagon + cell population. Pancreatic sections were obtained from eleven patients undergoing surgery for either a malignancy or pancreatitis (B,C,E-G, circle or square data points, respectively). Seven patients were non-diabetic (ND) and four had diagnosed type 2 diabetes (T2D). Statistical significance for the data was determined using a paired Student’s t-test. *, P < 0.05. Error bars indicate SEM.
cadaveric donor data (Figure 4D), we did not observe any increase in the percentage of GLP-1+/glucagon+/cell subpopulation in islets from donors with type 2 diabetes (Figure 5f).

4. DISCUSSION

The existence of intra-islet GLP-1 within rodent and human islets has been previously documented [2,11,12], however the presence of GLP-1 within islets does not necessarily provide evidence for any functional role. Despite several recent studies using genetic mouse models providing support for a functional role of intra-islet GLP-1 [15,16], the concept of a paracrine role for GLP-1 within human islets remains controversial and requires more evidence obtained from live tissue. As the GLP-1 antibody specificity is central to our methods and data interpretation, we have performed extensive in-house validation of the antibodies to confirm the following: 1) the active GLP-1 assay used detects only the active GLP-1 (7–36) isoform and not the 1–37, 9–37 inactive GLP-1 isoforms or glucagon (Figure 1L); and 2) the GLP-1 amidated C-terminal antibody used for quantification of the GLP-1 expressing a cell subpopulation does not cross-react with glucagon (Figure 2F).

Our results corroborate previous findings [23] that isolated human islets secrete robust amounts of active GLP-1 and that GLP-1 is likely originating from a significant subpopulation of a cells. Furthermore, the key prohormone convertase PC1/3 required for the proglucagon processing to release GLP-1 [10] is significantly increased in the GLP-1+/a cell subpopulation. Importantly, we observed an increase in the GLP-1+/a cell population in T2D islets, suggesting that metabolic factors such as hyperglycemia and chemokine action may regulate intra-islet GLP-1 expression. Indeed, hyperglycemia is associated with a cell PC1/3 expression in diabetic rodent models [23,24]. β cell injury has been shown to increase GLP-1 expression in human islets that involves the expression and secretion of the chemokine, SDF-1α [21], whereby β cells secrete SDF-1α that acts upon a cells to increase PC1/3 expression and GLP-1 production. Furthermore, the cytokine IL-6, produced from skeletal muscle and adipose tissue, has been shown to function as a metabolic signal that triggers islet a cells to increase islet GLP-1 production [22].

The possibility remains that intra-islet GLP-1 expression is increased as a result of the use of genetically modified models and/or the peri- and post-isolation procedure of human islets that may result in the β cell injury described above. However, we observed the presence of a significant subpopulation of GLP-1+/a cells in islets from whole pancreas sections that did not undergo any isolation stress, as well as in pancreatic sections obtained from living donors undergoing surgery [25,26]. Furthermore, we observed a significant increase in the number of GLP-1+/a cells in islets from T2D cadaveric donors when compared with ND islets, despite the isolation procedure and culturing conditions being identical. It should be noted that we did not observe a similar finding in islet sections from living donors. One possible explanation for this discrepancy is that all the donors undergoing surgery had pre-existing pancreatic disease (malignancy or pancreatitis) that may have masked any differences between donors with or without type 2 diabetes. We also acknowledge that a potential limitation of our study is the lack of information on any anti-diabetic medications being taken by the donors that might affect the results of our study. In conjunction, our results provide additional evidence for a potential physiological role for GLP-1 in human islets rather than an epiphenomenon resulting from isolation and culturing. The recent observation also supports the concept that active GLP-1 can be detected in human pancreatic tissue extracts at levels that were higher than observed in mouse pancreases [27].

We observed a strong GLP-1 secretory phenotype coupled with a high percentage of GLP-1+/a cells in human islets, therefore the potential for a functional paracrine role for GLP-1R signalling in human islets is of importance. Using the GLP-1R antagonist exendin-9, we have demonstrated that GLP-1R signalling plays a significant role in modulating intracellular calcium and GSIS and that islets from T2D donors are dependent on GLP-1R activation for >60% of insulin secretion. This observation is further supported by a significant increase in GLP-1+/a cells in T2D islets. Moreover, the effect of exendin-9 on insulin secretion at low glucose levels in T2D islets is more pronounced in T2D islets and may partially explain the higher levels of insulin secretion at low glucose observed in T2D islets. Because exendin-9 is a specific antagonist of GLP-1Rs, a plausible explanation is that exendin-9 is competitively inhibiting the actions of intra-islet GLP-1 on β cell GLP-1Rs [28], although an alternative explanation may be that glucagon also directly activates GLP-1Rs to stimulate insulin secretion. Additionally, a recent study that used gcr-/− and GLP-1R-/− genetic mouse models suggests that intra-islet GLP-1 levels are very low in wild-type mice and that glucagon can regulate insulin secretion via GLP-1R signalling, albeit at lower potency than GLP-1 [29]. However, previous reports indicate that glucagon acts via the glucagon receptor [30,31]. Moreover, GLP-1 is a ~400-fold more potent agonist at the GLP-1R than glucagon [32]. In this study we determined that there are large inherent differences in both the levels of intra-islet GLP-1 secretion and a cell population density between human and mouse islets that further highlight the controversy surrounding a paracrine role for GLP-1 in human physiology if only mouse models are used. Furthermore, due to the inherent phenotypic variability of human islets when compared to mice, this variation constitutes a limitation of our study. However, we have recently analysed GLP-1 secretion from human islets in association with islet isolation and donor parameters that attempts to characterise this potential limitation [33]. In the absence of difficult to obtain in vivo human data to further support the concept of intra-islet GLP-1, our study provides additional evidence for a paracrine GLP-1R signalling axis in human islets, perhaps via the localized high levels of GLP-1 secretion observed in this study. Future studies that quantify GLP-1R protein expression in the cell membranes of β cells of ND and T2D islets will help to establish if the increased GLP-1 expression we observe in the α cells of T2D islets is associated with an increase in its canonical receptor on β cells. However, in light of recent findings from mouse and human islets, a direct role for a cell derived glucagon acting upon β cell GLP-1Rs should also be considered [34,35].

The role for DPP4 and the clinically used DPP4 inhibitors on this intra-islet GLP-1 axis is also of interest. We tested the effects of the DPP4 inhibitor sitagliptin to evaluate whether some of the clinical efficacy of this class of drugs can be attributed to a direct intra-islet effect. Our flow cytometry analysis showed that DPP4 expression is relatively restricted to a cells, arguing for a regulatory role for DPP4 of a cell substrates such as GLP-1. As previously shown [4,36,37], we were also able to increase active GLP-1 in long-term human islet cultures. However, short-term perfusion of human islets with sitagliptin did not significantly increase GSIS in either ND or T2D islets; a result that is in direct contrast to previous human islet studies [36,37]. This discrepancy may be a result of various isolation, culture, and experimental conditions among research groups. Furthermore, we cannot exclude the possibility that intra-islet glucagon levels might contribute significantly to, or perhaps even dominate, activation of the GLP-1Rs in our
perfusion experiments [34,35,38], thus masking any enhancement in GSIS by increased levels of active GLP-1. Finally, DPP4 inhibitors may also improve islet function and survival and therefore indirectly enhance β cell function and insulin secretion [36,37].

In conclusion, our results provide evidence for the robust secretion of active GLP-1 from a subpopulation of α cells and an important paracrine role for GLP-1R signalling within human islets. The α-cell subpopulation is increased in T2D and is associated with a greater dependency on GLP-1R signalling for insulin secretion, suggesting that the α and β cells within human islets have adapted in T2D to amplify the paracrine pathway in an attempt to support insulin secretion.

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CONFLICT OF INTEREST

None declared.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data to this article can be found online at https://doi.org/10.1016/j.molmet.2020.101014.

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