Intravital imaging of islet Ca²⁺ dynamics reveals enhanced β cell connectivity after bariatric surgery in mice

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Bariatric surgery improves both insulin sensitivity and secretion and can induce diabetes remission. However, the mechanisms and time courses of these changes, particularly the impact on β cell function, are difficult to monitor directly. In this study, we investigated the effect of Vertical Sleeve Gastrectomy (VSG) on β cell function in vivo by imaging Ca²⁺ dynamics in islets engrafted into the anterior eye chamber. Mirroring its clinical utility, VSG in mice results in significantly improved glucose tolerance, and enhanced insulin secretion. We reveal that these benefits are underpinned by augmented β cell function and coordinated activity across the islet. These effects involve changes in circulating GLP-1 levels which may act both directly and indirectly on the β cell, in the latter case through changes in body weight. Thus, bariatric surgery leads to time-dependent increases in β cell function and intra-islet connectivity which are likely to contribute to diabetes remission.

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n estimated 415 million individuals in the world are living with diabetes, or ~1 in 11 adults1, with Type 2 diabetes (T2D) being the most prevalent form. In the United Kingdom it is predicted that by 2025 more than five million people will be diagnosed with the disease2. In response to this epidemic, an abundance of pharmacological and behavioural interventions have been utilised but often focus on glycaemic control rather than long-term disease resolution3,4. Several clinical trials have now reported that bariatric surgery, a group of gastrointestinal procedures originally developed to aid weight loss, can induce long-term remission of diabetes, with patients often achieving normoglycaemia without the need for any more glucose-lowering medications5–7.

Numerous studies8–12 have attempted to unravel the mechanisms by which blood glucose control is improved post-operatively. One hypothesis implicates augmented post prandial release of gastro-intestinal incretin hormones, particularly glucagon-like peptide 1 (GLP-1), which enhances glucose-stimulated insulin secretion13. This approach was established by Berggren and colleagues27 and has recently been developed by ourselves26 to assess coordinated islet β-cell function in living humans or preclinical models, that can often initiate Ca2+ waves (sometimes referred to as “hubs”), underlie the loss of insulin secretion observed in response to challenges associated with diabetes such as glucolipotoxicity23,25,26. Untangling these functional changes from alterations in β-cell mass in vivo is challenging, since the latter can only reliably be determined post mortem via pancreatic biopsies, and thus at a single time point.

In an effort to overcome these limitations, the present study aimed to directly and longitudinally observe the effect of vertical sleeve gastrectomy (VSG) on pancreatic β-cell function in mice, by transplanting “reporter” islets in the anterior chamber of the eye. This approach was established by Berggren and colleagues27 and has recently been developed by ourselves26 to assess coordinated islet behaviour in vivo. Importantly, this technique has allowed us to image Ca2+ dynamics recursively, in the same islet, over time and with near single-cell resolution, following surgery. We show that VSG increases β-cell Ca2+ dynamics within 8 weeks post-surgery when compared to pre-operative baseline and a sham-operated group. Moreover, we demonstrate that VSG increases the number and strength of β to β-cell connections at 10 weeks after surgery. These changes were associated with increased circulating GLP-1 levels, suggesting that enhanced incretin production contributes to postoperative improvements in β-cell performance.

**Results**

**VSG improves glucose tolerance.** Our experimental protocol is summarised in Fig. 1a. In brief, mice were placed on a high-fat

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**0.01 at 30, 60 and 90 min) as assessed by oral glucose tolerance test (OGTT) at week 2 (Fig. 1c) and intraperitoneal glucose tolerance test (IPGTT) 4 and 10 weeks post-operatively (p < 0.05 at min 30 and p < 0.001 at 60 and 90 min, Fig. 1d and 1e respectively). Strikingly, in all tolerance tests performed on VSG-treated mice, glucose peaked at 15 min post-glucose injection (3 g/kg) and dropped to baseline levels within 60 min by week 8, and near baseline levels at week 10. In contrast, in sham-operated mice, glucose peaked at 30 min and did not fully recover within the first 2 h of measurement.

**VSG improves insulin secretion and sensitivity but does not increase β-cell mass.** In order to understand the marked increase in the rate of glucose clearance in mice that had undergone VSG, we measured insulin secretion in vivo as a response to an IP glucose load (3 g/kg). Insulin secretion was increased significantly in VSG versus sham mice as early as 4 weeks post-operatively (Fig. 1f), with the observed peak at 15 min almost threefold higher compared to sham mice (p < 0.05). VSG mice were also significantly more insulin sensitive when compared to sham mice, as assessed by intraperitoneal insulin tolerance test at week 8 (ITT, 1.5 U/kg) (p < 0.001, Fig. 1g). However, pancreatic β-cell mass was not increased in the VSG group relative to sham controls at week 10 post-operatively (Supplementary Figs. 1a and 2). Notably, the ratio of α to β-cell mass was significantly higher in the VSG group, yet α-cell mass was not significantly increased (Supplementary Figs. 1b, c and 2).

**VSG enhances GLP-1 secretion.** To assess whether enhanced incretin release may contribute to the glucose-lowering effect of VSG we observed during IPGTT and OGTT, we measured plasma GLP-1 levels during fasting and 15 min following an orally administered glucose load (3 g/kg) (Fig. 1i). The concentration of GLP-1 was significantly higher in the VSG group, when compared to sham, 15 min post gavage (Fig. 1h).

**β-Cell Ca2+ dynamics are enhanced following VSG.** In order to explore changes in β-cell function after surgery, we monitored intracellular Ca2+ changes prospectively and in the same islets by confocal imaging of the anterior eye chamber26,27. Ca2+ increases, measured at ambient blood glucose concentrations in the range 12.5 ± 0.7 mmol/L for both VSG-treated and sham mice, which occurred at a single or multiple site across the islet but did not advance across the islet, were defined as “oscillations” and were categorised as level 1 (Supplementary Movie 2). Increases that had a defined site of origin and spread across a few cells but not the full width of the imaged plane were defined as “partial” waves (Fig. 2bi and Supplementary Movie 4) and were categorised as level 2. Those increases spreading across the whole islet were termed “waves” (Fig. 2ai, bii and Supplementary Movies 1 and 5) and were categorised as level 3. If the latter wave type was recurrent, we defined the behaviour as a “superwave” (Fig. 2biii and Supplementary Movie 6) and this was categorised as level 4. As illustrated in Fig. 3a, when imaged 0, 4 and 10 weeks after surgery, islets in sham-operated animals displayed a progressive
loss of coordinated Ca\textsuperscript{2+} dynamics, as defined by the proportion of the different wave types. Thus, when imaged at 0 weeks (Fig. 2ai), wave behaviour (beginning at the bottom right; red area) moved rapidly across the areas identified in yellow and blue. Comparable behaviour was seen at 4 weeks, with a similar site of origin of the wave (Fig. 2aii) but was lost at 10 weeks post sham surgery even though there was no significant weight difference between the two groups (Fig. 2aiii and Supplementary Movie 3).

In contrast, islets implanted into mice subject to VSG displayed sustained or gradually improving Ca\textsuperscript{2+} dynamics following surgery. Thus, the islet shown in Fig. 2bi initially showed partial wave activity but progressed to full wave activity by week 4 (Fig. 2Bii and Supplementary Movie 4) and to superwave by week 10 (Fig. 2biii and Supplementary Movie 6). Remarkably, almost all islets transplanted into VSG animals displayed either wave or superwave behaviour by week 8, even if VSG-treated animals did not display further weight loss. This is significantly higher when
Fig. 1 VSG improves glucose and insulin tolerance in HFHSD mice. a Timeline of procedures. b Body weight monitoring following VSG (red, n = 6 until week 6, then n = 4 animals) or sham surgery (black, n = 6 animals) (week 1: p = 0.006, week 2: p = 0.008, week 3: p = 0.02, week 4: p = 0.02, week 5: p = 0.009, week 6: p = 0.03). c Glucose was administered via oral gavage (3 g/kg) after mice were fasted overnight and blood glucose levels measured at 0, 15, 30, 60 and 90 min post gavage, 6 weeks after surgery, n = 5–6 mice/group. d Glucose was administered via intraperitoneal injection (3 g/kg) after mice were fasted overnight and blood glucose levels measured at 0, 15, 30, 60 and 90 min post injection, 4 weeks after surgery, n = 6 mice/group. p = 0.039 at 30 min, p < 0.001 for other timepoints. e Glucose was administered via intraperitoneal injection (3 g/kg) after mice were fasted overnight and blood glucose levels measured at 0, 15, 30, 60 and 90 min post injection, 10 weeks after surgery, n = 4–5 mice/group. f Corresponding insulin secretion levels measured on plasma samples obtained during the OGTT performed in d (n = 6 animals), p = 0.015 at 15 min, p = 0.027 at 30 min. g Insulin was administered via intraperitoneal injection (1.5 IU/kg) after mice were fasted for 5 h and blood glucose levels measured at 0, 15, 30, 60 and 90 min post injection, 7–8 weeks after surgery, n = 6 mice/group. h Corresponding GLP-1 secretion levels measured on plasma samples obtained during the OGTT performed in c (n = 9–10). I Corresponding glucose levels for 0, 15 min obtained during the OGTT performed in c (n = 10–14 animals), p = 0.0006. *p < 0.05, **p < 0.01, ***p < 0.001 VSG vs. Sham, following two-sided Student’s t-test or two-way ANOVA (adjusted for multiple comparisons). Data are expressed as means ± SEM. VSG vertical sleeve gastrectomy, HFHSD high-fat, high-sugar diet, IPGTT intraperitoneal glucose tolerance test, ITT insulin tolerance test, OGTT oral glucose tolerance test, GLP-1 glucagon-like peptide 1. Source data are provided as a Source Data file.

Compared to sham mice at the same time point (p = 0.02) (Fig. 3a). By week 10, the activity of all sham-transplanted islets dropped to almost zero (p = 0.004) (Fig. 3a). Mean wave front velocity, a measure of the speed of the wave calculated by distance (µm) divided by time (s), across the islet was not different between groups at any time point explored. Similarly, no differences were apparent between wave velocities for the different wave types in either VSG or sham operated (Fig. 3b).

VSG maintains the number and strength of β-cell–β-cell connections. Coordinated activity of β-cells is a feature of the healthy islet, and is likely to be important for the regulation of pulsatile insulin secretion29. As shown in Fig. 4a, b, Pearson-correlation analysis revealed no differences in apparent connectivity at week 0 (prior to surgery), whereas a progressive decline in connectivity was observed in the sham group. The number of connected cells (Fig. 4b) or the mean connectivity strength (R) (Fig. 4c, d) remained relatively constant in the VSG group, such that by week 10 these islets displayed significantly greater connectivity than the sham group (Fig. 4b). In summary, glucose-related Ca²⁺ signaling in VSG mice was characterised by higher magnitude and higher sensitivity to glucose when compared with sham mice or to pre-operative levels. Conversely, the sham group which continued to gain weight and experience further deterioration in their glucose tolerance revealed a diminution in coordinated islet behaviour suggestive of glucolipotoxic effects on islet function.

Liraglutide treatment does not fully replicate the Ca²⁺ activity observed in VSG. We sought next to test the impact of pharmacological GLP-1R agonism on islet Ca²⁺ dynamics in the absence of surgery. To do so, we treated obese, diabetic mice maintained on a high-fat diet (HFD) with the pharmacological agonist liraglutide daily for 6 weeks, and used a food-restricted, weight-matched control group. Although both groups lost a similar amount of weight during the treatment period (Supplementary Fig 3a), the liraglutide group displayed significantly improved glucose tolerance versus the pair-fed group from treatment week 2 to week 6 (Supplementary Fig. 3b, c). However, islet Ca²⁺ activity (Supplementary Fig. 3d), and subsequently the number of connected cells (Supplementary Fig. 3e), and the mean connectivity strength (R) (Supplementary Fig. 3f) were not different between the groups. In summary, additional effects of GLP-1R agonism on islet Ca²⁺ dynamics, beyond those of weight loss, were not apparent.

Mitochondrial membrane potential is not altered in ACE islets following VSG. To further explore the mechanism(s) involved in the observed changes in islet Ca²⁺ dynamics after VSG, we attempted an indirect measurement of glucose metabolism (and thus signalling) in β-cells within the engrafted islets, using the mitochondrial membrane potential sensor rhodamine 123 (TMRM). However, we did not observe any significant differences in steady-state mitochondrial membrane potential in islets from VSG versus sham-operated animals (Supplementary Fig. 2).

GLP-1R antagonism partly blunts the improvement in β-cell Ca²⁺ dynamics after VSG. In order to shed light on the possible mechanism(s) behind the enhanced postoperative β-cell Ca²⁺ activity, we repeated the study described above, exploring the effects of a GLP-1 receptor antagonist (Fig. 5a). From week 10, all animals (VSG and sham-operated) were administered the GLP-1 receptor antagonist Exendin (9-39) (Exendin9), via a subcutaneous osmotic pump at a rate of 10 pmol/kg/min, for 2 weeks. Ca²⁺ increases were monitored in the islets present in the anterior eye chamber by confocal imaging, as above, and ambient blood glucose concentrations were in the range 14.1 ± 0.5 mmol/L for both groups.

Following Exendin9 administration, a non-significant drop was observed in the activity (wave categories) of islets from the VSG group at week 12 (post-Exendin9) versus week 8 (pre-treatment), while in contrast a gradual increase in activity as assessed using this parameter continued in the sham group (Fig. 5b, c). A similar observation was made for the number of connected cells (Fig. 5d, e), and the mean connectivity strength (R) in VSG islets treated with Exendin9, when compared to week 8 (Fig. 5f–g) using baseline correction (i.e. normalisation to activity at week 0). These findings suggest that changes in circulating GLP-1 levels may contribute to the altered islet Ca²⁺ dynamics observed after VSG.

Plasma lipidomic profile is altered following VSG. We applied a targeted -omics approach to study small polar metabolite and lipid concentrations in peripheral plasma samples from sham and VSG-treated mice (Supplementary Fig. 4). Of 29 polar metabolites assessed, the levels of none were significantly altered in VSG animals when compared to sham-operated animals (Supplementary Fig. 4a).

In the lipidomic analysis, 298 lipid species from 17 different classes were studied (Supplementary Fig. 4b). Of these, one lipid from the glycerolipids class and two phosphatidylcholines, known for their role in membrane-mediated cell signalling30, were significantly upregulated in the VSG group (p < 0.01).

Discussion

Using an intravital imaging approach developed in recent years to monitor islet function in vivo26,27, we provide here evidence that VSG causes a dramatic improvement in β-cell Ca²⁺ connectivity,
Fig. 2 Description of Ins1Cre:GCaMP$f/f$ islet Ca$^{2+}$ dynamics: superwave, full wave, partial wave and no activity. Ins1Cre:GCaMP$f/f$ islets implanted in the anterior chamber of the eye and imaged for 400 frames (133 s) using a spinning disk confocal microscope (see “Methods”) at baseline week 0, postoperative week 4 and week 10. Each islet is separated in three regions of interest (ROIs) in order to categorise its activity. Red represents area 1 (distal islet), yellow represents area 2 (middle islet) and blue represents area 3 (proximal islet). Mean intensity is measured in each ROI for each frame (3 frames/s) and presented as Ca$^{2+}$ dynamics.

a. Ins1Cre:GCaMP$f/f$ islets implanted in a sham animal and imaged at (i) week 0 (full wave), (ii) week 4 (full wave) and (iii) week 10 (inactive).

b. Ins1Cre:GCaMP$f/f$ islet implanted in a VSG-treated animal (i) at week 0 (partial wave), (ii) week 4 (full wave) and (iii) week 10 (superwave). Scale: 100 μm. Plasma glucose levels during imaging were 12.5 ± 0.7 mmol/L.
postoperative follow-up and, by week 8, there was no significant weight difference between the sham and VSG groups. This allowed us to separate marked improvements in insulin secretion from significant weight loss without the need to pair-feed the sham group. Moreover, it corresponds with our previous findings in lean VSG-treated mice that demonstrated no weight difference when compared to sham mice at 4 weeks post-operatively, yet displayed improved glucose tolerance and corresponding insulin secretion curves during an IPGTT. Of note, insulin tolerance tests at week 8 demonstrated that VSG mice had improved insulin sensitivity, an effect previously attributed in humans to rapid and significant enhancement of postoperative hepatic insulin clearance.

Improved glucose tolerance in VSG-treated animals was underpinned by increased insulin secretion in response to glucose at 15 and 30 min, consistent with previous studies using VSG models. The fact that the insulin response to IPGTT was equally robust suggests that this effect is not solely due to an increased spike in blood glucose associated with elevated gastric emptying rates and upregulated glucose absorption, as has been previously postulated. Increased insulin secretion in the face of lower plasma glucose demonstrates enhanced β-cell glucose sensitivity, consistent with cell-autonomous changes in islet function, alterations in circulating levels of other regulators of secretion, or an increase in β-cell number. Analysis of the endogenous pancreatic β-cell mass showed no increase in VSG versus sham-operated mice, pointing to a functional change rather than a change in endogenous β-cell mass, as underlying increased insulin output. Moreover, and though this could not be quantified accurately due to the lack of focal distances stacking data, we saw no evidence for a change in the β-cell mass of islets engrafted into the eye. This is in line with previous findings demonstrating that there is no islet hyperplasia or increased β-cell turnover following bariatric surgery in humans or rats with obesity. These findings contrast other studies that have reported—over similar time-scales—increasing β-cell mass, or decreasing β-cell mass differences, which may reflect preoperative metabolic state or other factors.

Given many reports of increased GLP-1 release after bariatric surgery in both humans and rodents, here we explored levels of both circulating fasting and post-glucose gavage GLP-1 levels. We demonstrated that GLP-1 secretion was significantly higher in the VSG group 15 min after oral glucose gavage versus sham-operated animals (Fig. 1h). Apart from increasing glucose-stimulated insulin secretion and enhancing insulin gene transcription, GLP-1 also inhibits β-cell function and proliferation in rodent models of diabetes. Thus, although the underlying mechanisms remain unclear, increased GLP-1 levels might provide a partial explanation for the enhanced responses to glucose and the euglycemic effects observed after surgery. A number of studies have shown that, in postoperative patients with T2D, GLP-1 receptor (GLP-1R) blockade with the GLP-1R antagonist Exenclins causes significant reduction of insulin secretion when compared to a control group with lower GLP-1 levels, indicating an effect on β-cell function. Nonetheless, other studies have demonstrated that blocking GLP-1R in bariatric patients impairs glucose tolerance but not to a greater degree than before surgery, or when compared to non-operated patients. Furthermore, Ye and colleagues found that pharmacological or genetic blockade or elimination of GLP-1R signalling in rats or mice, respectively, had no impact on the ability of RYGB to lower body weight.

In our study, we administered Exenclins during postoperative weeks 10–12 (when the most marked difference in islet Ca2+ activity between groups was observed in our first cohort) via a subcutaneous osmotic pump, in order to assess if GLP-1R antagonism will blunt...
the effects on Ca\textsuperscript{2+} signalling. In line with the view that actions of GLP-1 may be involved in the improved islet Ca\textsuperscript{2+} dynamics, after 2 weeks of treatment Exendin9 partly reversed the differences in wave categories observed between VSG and sham groups (Fig. 5).

However, we do not exclude the possibility that a small change in body weight—which fell below significance after 8 weeks (Fig. 1b)—as well as alterations in insulin sensitivity (Fig. 1g), may also be involved in the change in β-cell function. While our -omics analysis also pointed towards an altered lipidomic profile (Supplementary Fig. 4b), the sample size was limited, restricting the conclusions that can be drawn.

Of note, differences in weight between the two groups (VSG and sham) were apparent after Exendin9 treatment (Fig. 5a). In line with this conclusion, administration of the GLP-1R agonist
liraglutide to HFD-fed animals which had not undergone surgery failed to fully recapitulate the alterations in islet Ca\(^{2+}\) dynamics when animals were carefully weight matched by pair feeding (Supplementary Fig. 3).

An important aspect of the present study has been to examine, at the cellular level, the functional connectivity between β-cells before and after VSG or sham surgery. The percentage of significantly connected cell pairs and correlation coefficient decreased substantially in the sham group at week 10, while in the VSG group these parameters remained stable for the duration of the study. We would note that hub/follower behaviour (i.e. the existence of a “power law” in the degree of connectedness)\(^2\) was not readily apparent in the present study. More rapid acquisition rates are likely to be needed to reveal such a hierarchy. Furthermore, we note that wave-like behaviour is more often apparent in the islet in vivo using GCaMP6f as the Ca\(^{2+}\) sensor\(^2\) than in some of our own and others’ earlier studies\(^2,23\) using entrapped, synthetic Ca\(^{2+}\) probes.

Changes in metabolic signalling by glucose, particularly changes in the glycolytic and oxidative metabolism of the sugar by mitochondria, are observed in islets from human subjects with T2D\(^8\) and are implicated in the recovery of glucose tolerance after Roux-en-Y-like gastric surgery in the mouse.\(^69\) In the former case, this is likely to involve decreases in the expression of the critical β-cell glucose transporter, GLUT2/SLC2A2, and in the low affinity glucose phosphorylating enzyme glucokinase (GCK).\(^9\) Similar observations have been made in a rodent T2D model, the diabetic db/db mouse.\(^7\) Measurements of gene expression were technically challenging in the current system as would have required post hoc surgical removal of islets from the eye, risking mRNA degradation and contamination with non-islet material. We instead attempted an indirect measurement of glucose metabolism, using the mitochondrial membrane potential sensor TMRM to measure this parameter in vivo. Our data with this approach did not, however, provide compelling evidence for such changes after surgery, though our sample size was relatively small. It is conceivable, therefore, that changes in the expression of genes more directly associated with the control of Ca\(^{2+}\) dynamics and connectivity in and between β-cells, including Trpm2 and Gjd2 (Cx36)\(^60\) and others associated with insulin secretion\(^4\), play a role in the improvements observed post-surgery.

Limitations of the study. Although the surgical model used provides us with original information on β-cell activity via continuous monitoring, there are undoubtedly limitations in the use of islets engrafted into the ACE. These include potential differences between the vascularisation and innervation at this site compared to pancreatic in situ islets\(^12\). Our findings on ACE-engrafted islet reactivation following VSG are however in line with previous results focusing on pancreatic islets isolated post mortem. Thus, Douros et al.\(^4\) performed Ca\(^{2+}\) imaging in vitro in mouse islets post mortem and showed that the percentage of islets displaying Ca\(^{2+}\) oscillations in response to glucose was enhanced 2.2-fold in the VSG group, indicating increased islet glucose sensitivity after surgery. However, these earlier studies were cross-sectional in nature, and as such did not explore the apparent reactivation in vivo of individual islets and islet subdomains, in the living animal, as described here. Finally, we note that the use of isoflurane as an anaesthetic agent has recently been linked to glucose increases\(^7\) and might therefore have an impact on Ca\(^{2+}\) measurements. In our studies, the average fed glucose was constant between VSG and sham groups at an average 13.3 ± 0.5 mmol/L for both cohorts, and all animals were imaged at the same isoflurane dosage, yet the observed Ca\(^{2+}\) dynamics were markedly different between groups.

In conclusion, our findings indicate that bariatric surgery improves glycaemic control at least partially by maintaining: (a) β-cell function and (b) coordinated activity across the islet. These effects involve changes in circulating GLP-1 levels which may act both directly and indirectly on the β-cell, in the latter case through changes in body weight. Future challenges are to understand more fully the mechanisms through which these changes are effected at the paracrine, endocrine and cellular levels.

Methods

Animals. All animal procedures undertaken were approved by the British Home Office under the UK Animal (Scientific Procedures) Act 1986 (Project License PPL PA031F707 to LL) with approval from the local ethical committee (Animal Welfare and Ethics Review Board, AWERB), at the Central Biological Services (CBS) unit at the Hammersmith Campus of Imperial College London.

Adult male C57BL/6 mice (Envigo, Huntingdon, UK) were maintained under controlled temperature (21–23 °C), humidity (45–50%) and light (12:12 h light–dark schedule, lights on at 0700 hours) and were health screened clean for Federation of European Laboratory Animal Science Associations (FELASA) list pathogens. From the age of 8 weeks they were put on a 58 kcal% fat and sucrose diet (D12331, Research Diet, New Brunswick, NJ) ad libitum to induce obesity and diabetes. Four weeks after the start of this diet, the animals underwent islet transplantation into the anterior chamber of the eye of genetically modified islets expressing GCaMP6f to allow for intravital measurements of cytosolic Ca\(^{2+}\). Four weeks after islet transplantation, the animals underwent either a VSG or a sham surgery as described below. The number of the study was initially calculated using 10-week body weight (Sham: 34.5 ± 4.3, VSG: 29.3 ± 2.5 g) and 4-week AUC glycaemia values (Sham:1,467 ± 76 mmol/L, VSG: 1,181 ± 72 mmol/L from Garbutt and Cummings\(^9\)). This was calculated as a continuous endpoint, two independent sample study groups, a (type 1 error probability) 0.05 and β (type II error probability) 90%.

Ins1CreGCaMP6f\(^6\) mice, used as donors for islet transplantation, were generated by crossing Ins1Cre mice (provided by J. Ferrer, this Department of Metabolism, Digestion and Reproduction) to mice that express GCaMP6f downstream of a LoxP-flanked STOP cassette (The Jackson Laboratory, stock no. 028665). Islets donated from either sex (8 males, 6 females) were used for transplantation.
Islet transplantation into the anterior chamber of the mouse eye (ACE).
Pancreatic islets were isolated by injecting collagenase-containing medium into the pancreatic duct, followed by separation from the exocrine tissue by gradient density and handpicking. The islets were cultured for 24 h in RPMI 1640 medium containing 11 mM glucose supplemented with 2 mM L-glutamine, 10% heat-inactivated foetal calf serum, 100 IU/mL penicillin, and 100 μg/mL streptomycin. For transplantation, 10–20 islets were aspirated with a 27-gauge blunt eye cannula (BeaverVisitec, UK) connected to a 100 μL Hamilton syringe (Hamilton) via 0.4-mm polyethylene tubing (Portex Limited). Prior to surgery, mice were anaesthetised with 2–4% isoflurane (Zoetis) and placed in a stereotactic frame to stabilise the head. The cornea was incised near the junction with the sclera, being careful not to damage the iris. Then, the blunt cannula, pre-loaded with islets, was inserted into the ACE and islets were expelled (average injection volume 20 μL for 10 islets). Carprofen (Bayer, UK) and eye ointment were administered post-surgery.

Vertical sleeve gastrectomy. Three days before bariatric or sham surgery, animals were exposed to liquid diet (20% dextrose) and remained on this diet for up to 4 days post-operatively. Following this, mice were returned to high-fat/high-sucrose diet until euthanasia and tissues harvested 10 weeks post bariatric surgery. Anaesthesia was induced and maintained with isoflurane (1.5–2%). A laparotomy incision was made, and the stomach was isolated outside the abdominal cavity. A simple continuous pattern of suture extending through the gastric wall and along both gastric walls was placed to ensure the main blood vessels were contained.
Approximately 60% of the stomach was removed, leaving a tubular remnant. The edges of the stomach were inverted and closed by placing two serosa only sutures, using Lembert pattern. The initial full thickness suture was subsequently removed. Sham surgeries were performed by isolating the stomach and performing a 1 mm puncture in all mice following a 3-day wash period post-Exendin9 administration. Blood was sampled from the tail vein at 0, 15, 30, 60 and 90 min after glucose administration. Blood glucose was measured with an automatic glucometer (Accuchek; Roche, Burgess Hill, UK).

Glucose tolerance tests. Mice were fasted overnight (total 16 h) and given free access to water. At 0900 hours, glucose (3 g/kg body weight) was administered via intraperitoneal injection or oral gavage. Blood was sampled from the tail vein at 0, 5, 15, 30, 60 and 90 min after glucose administration. Blood glucose was measured with an automatic glucometer (Accuchec; Roche, Burgess Hill, UK).

Insulin tolerance tests. Mice were fasted for 8 h and given free access to water. At 1500, human insulin (Actrapid, Novo Nordisk) (1.5 U/kg body weight) was administered via intraperitoneal injection. Blood was sampled from the tail vein at 0, 15, 30, 60 and 90 min after insulin administration. Blood glucose was measured with an automatic glucometer (Accuchec; Roche, Burgess Hill, UK).

Plasma insulin and GLP-1 measurement. To quantify circulating insulin and GLP-1 (1–37) levels, 100 μL of blood was collected from the tail vein into heparin-coated tubes (Sarstedt, Beaumont Leys, UK). Plasma was separated by sedimentation at 10,000 g for 10 min (4°C). Plasma insulin levels were measured in 5 μL aliquots and GLP-1 (1–37) levels were measured in 10 μL aliquots by ELISA kits from Crystal Chem (USA).

Osmotic pump implantation. ALZET mini-osmotic pump Model 1002 (DURECT Corporation, Cupertino, CA, United States) were utilised for continuous subcutaneous administration. The pump had a fill volume of 100 μL and a predicted pumping rate was 0.25 μL/H for 14 days. Exendin9 (Insight Biotechnology Limited, UK) was completely dissolved in 0.9% saline within 1 min and filled into the pump according to the manufacturers protocol. The final rate was approximately 10 pmol/kg/min. After filling, pumps were primed in 0.9% saline (10 mL) 37°C for at least 3 h according to the manufacturers protocol. Each mouse received a subcutaneous implantation of one osmotic pump, placed between the shoulder blades, on week 10 following VSG or sham procedures.

Mitochondrial membrane potential measurement. Imaging sessions were performed as described above, with the mouse held in a stereotactic frame and the eye gently retracted, with the animal maintained under 2–4% isoflurane anaesthesia. All imaging experiments were conducted using a spinning disk confocal microscope (Nikon Eclipse Ti, Crest spinning disk, ×20 water dipping 1.0 NA objective). The signal from GCaMP6f fluorophore (ex. 488 nm, em. 525 ± 25 nm) and TMRM (ex. 549 nm, em. 575 nm) was monitored by obtaining a time-series experiments for up to 20 min at a rate of 12.5 ± 0.7 Hz. t-Tests were recorded for three mice, with a mean blood glucose reading of 12.5 ± 0.7 mmol/L. In cohort 1 (control 1 and 2) (Exendin9), islets were continuously monitored, and the focus was manually adjusted to counteract movement. Animals were imaged 3 days prior to VSG (baseline) and then at 4, 8, and 10 weeks post-operatively.

In vivo Ca2+ imaging of Ins1Cre:GCaMP6f/β-fl islets in the ACE. A minimum of 4 weeks was allowed for full implantation of islets before imaging. Imaging sessions were performed as previously described26 with the mouse held in a stereotactic frame and the eye gently retracted, with the animal maintained under 2–4% isoflurane anaesthesia. All imaging experiments were conducted using a spinning disk confocal microscope (Nikon Eclipse Ti, Crest spinning disk, ×20 water dipping 1.0 NA objective). The signal from GCaMP6f fluorophore (ex. 488 nm, em. 525 ± 25 nm) was monitored in time-series experiments for up to 20 min at a rate of 12.5 ± 0.7 Hz. Mice were fasted for 8 h and given free access to water. At 1500, human insulin (Actrapid, Novo Nordisk) (1.5 U/kg body weight) was administered via intraperitoneal injection. Blood was sampled from the tail vein at 0, 15, 30, 60 and 90 min after glucose administration. Blood glucose was measured with an automatic glucometer (Accuchec; Roche, Burgess Hill, UK).

Liraglutide study. Adult male C57BL/6J mice (25–30 g) (Charles River, UK) were individually housed under controlled temperature (21–23°C) and light (12:12 h light-dark schedule, lights on at 0700 hours) with ad libitum access to standard chow and water. Syngeneic Ins1Cre:GCaMP6f islets were implanted into the anterior eye chamber of their eyes. Prior to imaging experiments, animals were put on HFD for approximately 8 weeks to induce obesity and glucose intolerance. The energy density of HFD was 5.21 kcal/g and composed of 20% kcal protein, 60% kcal fat and 20% kcal carbohydrate (Research diets; D12492). A glucose intolerant phenotype was confirmed via IPGTT (1 g/kg glucose bolus), prior to baseline recordings of Ins1Cre:GCaMP6f islets (ex. 488 nm; 300 ms exp. at 3 Hz). Blood glucose readings during the acquisitions averaged around 12.2(±0.9) mmol/L. Subsequently, animals were allocated to receive daily subcutaneous injections of either liraglutide (at 50 nmol/kg) or vehicle (saline + Zn2+2) combined with weight-matching. Injections were administered over a period of 6 weeks. At the end of the treatment regimen, IPGTT’s (1 g/kg glucose bolus) and a final recording of calcium dynamics of Ins1Cre:GCaMP6f islets were conducted (ex: 488 nm; 300 ms exp. at 3 Hz). During imaging experiments, blood glucose measurements were comparable for the two groups, averaging around 10.2(±1) versus 9.2(±0.2) mmol/L for liraglutide and weight-matched groups, respectively. This study was conducted under project licence PPL759462 (project licence holder Prof. Kevin Murphy).

Immunohistochemistry of pancreas sections. Isolated pancreata were isolated 10 weeks after surgery (age: 32 weeks) and were fixed in 10% (vol/vol) buffered formalin and embedded in paraffin wax within 24 h of removal. Slides (5 μm) were submerged sequentially in HistoClear (Sigma, UK) followed by washing in decreasing concentrations of ethanol to remove paraffin wax. Permeabilised pancreatic slices were blotted with ready-diluted anti-guinea pig insulin (Agriculture Technologies, USA) and anti-mouse glucagon (Sigma, UK) primary antibody (1:1000). Slides were visualised by subsequent incubation with Alexa Fluor 488 (1:1000) and 568-labelled donkey anti-guinea pig and anti-mouse antibody (1:1000). Samples were mounted on glass slides using VectashieldTM (Vector Laboratories, USA) containing DAPI. Images were captured on a Zeiss Axio Observer.Z1 motorised inverted widefield microscope fitted with a Hamamatsu Flash 4.0 Camera using a Plan-Apochromat 200/0.8 M27 air objective with Colibri2 LED illumination. Data acquisition was controlled with Zeiss LSM 880 software. Fluorescence quantification was achieved using ImageJ (https://imagej.nih.gov/ij/). Whole pancreas was used to quantify cell mass.

Metabolic and lipodimetric measurements. Plasma was collected via cardiac puncture in all mice following a 3-day wash period post-Exendin9 administration. Molecules from plasma were measured using two approaches at Steno Diabetes Center Copenhagen. One method focused on a panel of known metabolites.
previously associated with diabetes, diabetes complications and metabolic dysfunction, these molecules were fully quantified using targeted ultra-high-performance liquid chromatography coupled triple quadruple mass spectrometry (UHPLC-QqtOQ-MS/MS) as described earlier\textsuperscript{72}. The second method aimed to measure a broad array of lipid species. Lipidomic sample preparation followed the Folch procedure with minor adjustments\textsuperscript{96}. Lipids were measured with UHPLC coupled quadruple-time-of-flight mass spectrometry (UHPLC-QTOF-MS) in both positive and negative ionisation mode. Identification was done using MZmine (version 2.2.8) matching to an in-house library\textsuperscript{77}, finally peak areas were normalised to internal standards\textsuperscript{78}. Significance was tested by Student’s two-tailed t-test.

Statistical analysis. Data were analysed using GraphPad PRISM 9.0 software. Significance was tested using non-parametric, unpaired Student’s two-tailed t-tests with Bonferroni post-tests for multiple comparisons, or two-way ANOVA as indicated. P < 0.05 was considered significant and errors signify ±SEM.

Pearson (R)-based connectivity and correlation analyses. Correlation analyses in an imaged islet were performed between β-cell pairs and their extracted fluorescent Ca\textsuperscript{2+} traces over time in MATLAB using a custom-made script. β-Cell intensity was measured in 100 consecutive frames (30 s). Regions of Interest (18–40 per islet, depending on size) were selected with single or near single-cell resolution (i.e. 10–20 μm diameter) and captured approximately 95% of the fluorescence of the image plane. A noise reduction function (effectively a rolling average) was applied to smooth noisy data and minimise the effects of outliers. The data window size used to calculate the moving averages was set to 5% of the total data points collected during each capture. The correlation coefficient R between all possible (smoothed) β-cell pair combinations (excluding the autocorrelation) was assessed using Pearson’s correlation. The Cartesian co-ordinates of the imaged cells were then incorporated in the construction of connectivity line maps. Cell pairs were connected with a straight line, the colour of which represented the correlation strength and was assigned to a colour-coded light–dark ramp \((R = 0.1–0.25 [blue], 0.26–0.5 [green], R = 0.51–0.75 [yellow], R = 0.76–1.0 [red])\). Cells with the highest number of possible cell pair combinations are shown in red. Data are also displayed as heatmap matrices, indicating all possible β-cell pair connections on each axis (min = 0; max = 1). The positive R values (excluding the auto-correlated cells) and the percentage of cells that were connected to one another were averaged and compared between groups.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. The authors declare that all data supporting the findings of this study are available within the paper and its supplementary information files. Source data are provided with this paper.

Code availability. The custom-made script that supports the connectivity findings of this study are available in GitHub with the identifier ‘Connectivity-calcium_Pearson-correlation-coefficient-analysis’\textsuperscript{99}.

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Additional information

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