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Title

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Journal Nature Metabolism, 1(11)

ISSN

2522-5812

Authors

Menegaz, Danusa Hagan, D Walker Almaça, Joana <u>et al.</u>

Publication Date 2019-11-01

DOI

10.1038/s42255-019-0135-7

Peer reviewed



HHS Public Access

Author manuscript *Nat Metab.* Author manuscript; available in PMC 2020 May 19.

Published in final edited form as:

Nat Metab. 2019 November ; 1(11): 1110–1126. doi:10.1038/s42255-019-0135-7.

Mechanism and effects of pulsatile GABA secretion from cytosolic pools in the human beta cell

Danusa Menegaz^{1,16}, D. Walker Hagan^{2,16}, Joana Almaça¹, Chiara Cianciaruso³, Rayner Rodriguez-Diaz¹, Judith Molina¹, Robert M. Dolan², Matthew W. Becker², Petra C. Schwalie^{3,4}, Rita Nano⁵, Fanny Lebreton⁶, Chen Kang^{7,8}, Rajan Sah^{7,8}, Herbert Y. Gaisano⁹, Per-Olof Berggren^{10,11,12}, Steinunn Baekkeskov^{3,13,*}, Alejandro Caicedo^{1,10,14,15,*}, Edward A. Phelps^{2,3,*}

¹ Division of Endocrinology, Diabetes and Metabolism, Department of Medicine, University of Miami Miller School of Medicine, Miami, FL 33136, USA. ² J. Crayton Pruitt Family Department of Biomedical Engineering, University of Florida, Gainesville, FL 32611, USA. ³ Institute of Bioengineering, School of Life Sciences, École Polytechnique Fédérale de Lausanne, CH-1015 Lausanne, Switzerland. ^{4.}Swiss Institute of Bioinformatics, CH-1015 Lausanne, Switzerland. ⁵ Pancreatic Islet Processing Facility, Diabetes Research Institute, IRCCS San Raffaele Scientific Institute, 20132 Milan, Italy ⁶. Cell Isolation and Transplantation Center, Faculty of Medicine, Department of Surgery, Geneva University Hospitals and University of Geneva, CH-1205 Geneva, Switzerland ⁷ Center for Cardiovascular Research and Division of Cardiology, Department of Internal Medicine, Washington University School of Medicine, St Louis, MO, 63110, USA ^{8.}Department of Internal Medicine, Division of Cardiovascular Medicine, University of Iowa, Carver College of Medicine, Iowa City, IA, 52242, USA. ⁹. Department of Medicine, University of Toronto, Toronto, ON, Canada M5S 1A8. ¹⁰ Diabetes Research Institute, University of Miami Miller School of Medicine, Miami, FL 33136, USA. ^{11.}The Rolf Luft Research Center for Diabetes & Endocrinology, Karolinska Institutet, Stockholm, SE-17177, Sweden. ¹² Division of Integrative Biosciences and Biotechnology, WCU Program, University of Science and Technology, Pohang, 790-784 Korea. ^{13.}Departments of Medicine and Microbiology / Immunology, Diabetes Center, University of California San Francisco, San Francisco, CA, 94143, USA. ¹⁴ Department of

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^{*}Correspondence: ephelps@bme.ufl.edu (E.A.P.), sbaekkeskov@ucsf.edu (S.B.), and acaicedo@med.miami.edu (A.C.). AUTHOR CONTRIBUTIONS

E.A.P. and S.B. conceived and carried out subcellular studies of GABA-ergic components in islet cells. D.M. and A.C. conceived and identified GABA release from islets in pulses and pioneered the biosensor cell technique for analyzing the dynamics of islet GABA release. E.A.P. conceived and identified the role of VRAC and TauT in GABA release and uptake. D.W.H. and E.A.P. analyzed the genetic models for LRRC8A^{-/-} MIN6 cells, β c-LRRC8A^{-/-} murine islets, and knock-down LRRC8A-shRNA human islets; D.M., D.W.H., and E.A.P. performed experiments to detect GABA, taurine, and serotonin/insulin secretion; J.M. and J.A. performed hormone assay experiments and ELISAs; J.A. conducted NPY-pHluorin experiments to measure exocytosis; H.Y.G. generated adeno-NPY-pHluorin vectors; R.S. generated genetic models for *LRRC8A* knock-out MIN6 cells and *LRRC8A*^{II/II} murine islets; C.K. isolated and shipped *LRRC8A*^{II/II} murine islets; M.W.B. isolated rodent islets and performed Western blot analyses; C.C. prepared cultures of primary rat hippocampal neurons; P.C.S. performed bioinformatics analysis; R.N. and F.L. isolated human islets for research; E.A.P., D.M., D.W.H., J.A., C.C., R.M.D., and R.R.-D. collected, analyzed, and quantified immunohistochemical data; P-O.B. provided critical equipment, reagents, expertise, and support. D.M., D.W.H., S.B., A.C., and E.A.P. designed the study, analyzed data, and wrote the paper. All authors discussed the results and commented on the manuscript.

The authors declare no competing interests.

Physiology and Biophysics, Miller School of Medicine, University of Miami, Miami, FL 33136, USA. ^{15.}Program in Neuroscience, Miller School of Medicine, University of Miami, Miami, FL 33136, USA. ^{16.}These authors contributed equally

Abstract

Pancreatic beta cells synthesize and secrete the neurotransmitter γ -aminobutyric acid (GABA) as a paracrine and autocrine signal to help regulate hormone secretion and islet homeostasis. Islet GABA release has classically been described as a secretory vesicle-mediated event. Yet, a limitation of the hypothesized vesicular GABA release from islets is the lack of expression of a vesicular GABA transporter in beta cells. Consequentially, GABA accumulates in the cytosol. Here we provide evidence that the human beta cell effluxes GABA from a cytosolic pool in a pulsatile manner, imposing a synchronizing rhythm on pulsatile insulin secretion. The volume regulatory anion channel (VRAC), functionally encoded by LRRC8A or Swell1, is critical for pulsatile GABA secretion. GABA content in beta cells is depleted and secretion is disrupted in islets from type 1 and type 2 diabetic patients, suggesting that loss of GABA as a synchronizing signal for hormone output may correlate with diabetes pathogenesis.

INTRODUCTION

The neurotransmitter γ -aminobutyric acid (GABA) occurs at high concentrations in the inhibitory neurons of the central nervous system and the pancreatic islets of Langerhans¹. The physiological purpose of GABA in islets was initially proposed to be a paracrine signal released from islet beta cells to inhibit alpha cells^{2–4}. Recent evidence suggests that GABA also has strong protective and regenerative effects on the beta cells themselves⁵. GABA increases beta cell mass in rodent and grafted human islets^{6–11} and ameliorates diabetes in non-obese diabetic (NOD) mice¹². Additionally, long-term GABA treatment in diabetic mice prevents alpha-cell hyperplasia¹³ and promotes alpha cell trans-differentiation into beta cells^{14,15}, although this latter effect is now disputed^{16,17}. Immune cells possess receptors for GABA^{18,19} which suppresses cytokine secretion, inhibits proliferation, and tempers migration^{10,18,20}. GABA inhibits autoreactive T cell proliferation at the interstitial concentrations found in islets (0.1–10 μ M)^{21–23}. Together, this evidence implicates GABA as a potent trophic factor and suppressive immunomodulator in islets. It is conceivable that the loss of GABA may leave islet regions vulnerable to inflammation²⁰.

GABA is synthesized by the enzyme glutamic acid decarboxylase (GAD), which is expressed as two isoforms, GAD65 and GAD67. Human beta cells only express the GAD65 isoform²⁴, which is detected in the cytosol and anchored to the cytosolic face of Golgi and peripheral vesicle membranes by hydrophobic modifications including palmitoylations^{1,25}. Earlier low resolution imaging studies localized GAD and GABA to synaptic-like microvesicles in beta cells^{26–28}. More recently, GABA has been detected in insulin granules from which it is released upon stimulation with glucose to activate GABA_A receptors in beta cells^{29–32}. However, a substantial fraction of the GABA pool is independent of extracellular glucose concentration and yet contributes significantly to GABA signaling in the

islet^{31,33,34}. The source of this pool of GABA secretion appears to be the cytosol³⁵, but a mechanism linking cytosolic GABA to extracellular release has remained unidentified. In analogy to the role ambient GABA plays in the central nervous system³⁶, such release of GABA may be crucial for regulating islet cell excitability, coordinating cell activity throughout the islet, and producing the beneficial effects mentioned above.

Here, we have assessed how GABA is released from human beta cells. We compared GABA release from a predominantly cytosolic pool of intracellular GABA in beta cells with that of GABA contained in vesicular membrane compartments including synaptic-like microvesicles and the larger insulin secretory vesicles. We provide evidence that cytosolic GABA is released from human beta cells via volume regulatory anion channels (VRAC) in a pulsatile pattern that is independent of glucose concentration. Furthermore, the GABA-permissive taurine transporter (TauT) mediates uptake of interstitial GABA. Finally, we studied the impact of this non-vesicular GABA release on insulin secretion in human islets from non-diabetic and diabetic donors.

RESULTS

Cytosolic pools of GABA are depleted in type 1 and type 2 diabetic islets

Earlier studies have shown that GABA is present at high levels in pancreatic islets^{35,37}, but due to the use of glutaraldehyde fixation, the resolution and the ability to use multiple antibody labeling of these early images was limited. Using an antibody that does not require glutaraldehyde fixation, we studied the GABA content in human islets from non-diabetic and diabetic donors. Human pancreas sections from non-diabetic donors immunostained for GABA, insulin, and glucagon showed that GABA is highly concentrated in islets compared to the surrounding exocrine tissue (Figure 1a). GABA staining was strongest in beta and delta cells, while alpha cells contained little or no GABA (Figure 1a,b, Extended Data 1).

Comparing the GABA content in human pancreas sections from non-diabetic (Figure 1c), type 2 diabetic (Figure 1d), and type 1 diabetic (Figure 1e,f) donors, we observed that type 1 and type 2 diabetic islets were depleted of GABA (Figure 1g), a result, which has not been previously described. In type 1 diabetic islets, the loss of GABA was not only observed in islets devoid of beta cells. Rather, even islets with residual beta cells were depleted of GABA (Figure 1f). Taurine, a small molecule compound with molecular characteristics similar to GABA, was not depleted in diabetes, indicating that the loss of GABA is unlikely caused by varying sample quality or fixation (Extended Data 1). The loss of GABA in type 1 and type 2 diabetic islets observed by histology was confirmed by high-performance liquid chromatography (HPLC) measurements of the GABA content of isolated whole human islets from non-diabetic, type 1 diabetic, and type 2 diabetic donors (Figure 1h).

We next addressed whether the lack of GABA in beta cells from type 2 and/or type 1 diabetic patients was the consequence of a loss of expression of the GABA synthesizing enzyme GAD65 (Figure 1i,j). GAD65 immunoreactivity in beta cells of type 2 diabetic patients and in remaining beta cells in type 1 diabetes was similar to non-diabetic controls (Figure 1k–n). Thus, the lack of GABA in diabetic islets is not due to lack of GAD65 expression in beta cells.

In human beta cells, GAD65 was highly expressed in Golgi membranes, peripheral vesicle membranes, and the cytosol, while human alpha cells expressed GAD65 at low levels and the localization was restricted to ER/Golgi membranes while lacking in vesicle membranes and the cytosol (Extended Data 1). Remarkably, alpha cells contained little or no GABA (Figure 1a,b) despite expressing GAD65. Expression of GAD65 in alpha cells of diabetic pancreases was indistinguishable from non-diabetic pancreases.

Subcellular localization suggests a non-vesicular GABA release mechanism in beta cells

Despites its well-established importance to islet physiology, the dominant mechanism of GABA release from islets remains unclear. We searched for immunohistochemical evidence of the canonical synaptic-like microvesicle mechanism of GABA-release^{27,28}. In GABA-ergic neurons, hydrophobic post-translational modifications of the synthesizing enzyme GAD65 anchor this protein to the cytosolic face of synaptic vesicle membranes where it colocalizes with the vesicular GABA transporter (VGAT)^{38,39}. By association with GAD65, VGAT mediates transport of the product GABA into the synaptic vesicle lumen, where it accumulates in preparation for regulated secretion⁴⁰. The existence of an analogous GABA-secreting system in islet cells would require co-expression of GAD65 and VGAT⁴¹ or another as-yet unidentified vesicular GABA transporter in synaptic-like microvesicles. We found that almost all (> 99%) human beta cells lacked expression of VGAT. VGAT expression was, however, detected in a small subset of human beta cells as well as in the somatostatin producing delta cells, coinciding with the presence of GABA in vesicular compartments. (Figure 2a,b, Extended Data 2)

Human alpha cells were devoid of VGAT. In contrast, in rat islets, VGAT expression was mainly detected in alpha cells (Figure 2b, Extended Data 2) while predominantly absent in beta and delta cells (Supplementary Table 1). VGAT, also known as the vesicular inhibitory amino acid transporter (VIAAT), is a transporter for glycine in addition to GABA in neurons⁴⁰ and in rat alpha cells, where it localizes with glycine in secretory vesicles⁴². The absence of GABA in rat alpha cells in our analyses is consistent with VGAT transporting glycine but not GABA in those cells and is in contrast with the results in human delta cells, which express VGAT and also contain GABA. A table summarizing the cell type-specific expression of GABA, GAD65 and VGAT in human and rat islets is included as supplementary information (Supplementary Table 1).

To assess whether VGAT is required for accumulation of GABA in peripheral vesicles, we compared the subcellular localization of GABA and GAD65 with synaptic vesicle markers synaptophysin, SV2C, and VGAT in monolayers of human islet cells (Figure 2c–i) and in hippocampal neurons (Figure 2h,i, Extended Data 2). Almost all human beta cells (~99%) exhibited a uniform cytosolic staining pattern for GABA (Figure 2c). However, in the rare VGAT⁺ human beta cells (<1%; Figure 2d) or in human delta cells (Figure 2e), GABA exhibited a well-defined vesicular staining pattern that colocalized with VGAT. The VGAT⁺/GABA⁺ vesicular structures in beta cells colocalized with insulin, but not GAD65, identifying them as insulin secretory granules. The existence of a small population of VGAT ⁺ beta cells containing GABA in insulin granules reconciles our observations with previous reports of quantal exocytotic GABA release events that coincide with insulin release³¹.

These data notwithstanding, the majority of beta cells did not express VGAT and had cytosolic rather than vesicular GABA content.

While GABA itself is mainly cytosolic, the GABA-synthesizing enzyme GAD65 exhibits a strongly punctate staining pattern in beta cells (Figure 2f)¹. These GAD65 puncta in beta cells have been previously described as synaptic-like microvesicles^{27,28}. In human or rat beta cells containing predominantly cytosolic GABA, there was no concentration of GABA in GAD65-positive puncta or in insulin granules (Figure 2f–h). Furthermore, we could not identify any marker for synaptic vesicles or endosomes that strongly colocalized with GAD65 vesicles or GABA in human or rat beta cells (SV2C, synapsin 6, synaptophysin, syntaxin, VAMP2, VGAT, WIPI2, APPL1, caveolin, clathrin, EEA1, GOPC, LAMP1, LC3, Rab3c, Rab5, Rab6, Rab7, Rab8, Rab9, Rab10 and Rab11 were tested). Yet, synaptic vesicle markers show strong colocalization with both GAD65 and GABA in neurons (Figure 2h,i, Extended Data 2). Together, these results indicate that the magnitude of GABA release from beta cells does not involve synaptic-like microvesicles.

Islet GABA secretion is pulsatile, and depends on GABA content

To investigate GABA secretion from islets we used cellular biosensors^{43–45} (Figure 3 and Extended Data 3). GABA secretion in real time was monitored by recording intracellular Ca^{2+} mobilization ($[Ca^{2+}]_i$) in biosensor cells stably expressing heteromeric GABA_B receptors (GABA_B R1b and GABA_B R2) and the G-protein α subunit, $G\alpha qo 5^{46,47}$ (Extended Data 3). We found that GABA is secreted from human islets in rhythmic bursts (Figure 3a) independent of glucose concentration. Biosensor responses could be blocked by the selective GABA_B receptor antagonist CGP55845 (10 μ M) (Figure 3b) and did not occur in the absence of islets (Figure 3a), confirming that the $[Ca^{2+}]_i$ responses were elicited by GABA released from islet cells. Pulsatile GABA secretion had distinct periods that in most islet preparations ranged from 4 to 10 minutes (Figure 3c), similar to pulsatile insulin secretion in humans⁴⁸. The periodicity of GABA secretion varied little between islets from the same human islet preparation. Calibrating biosensor responses from islets by comparing them to responses evoked by direct application of GABA in the same experiment, GABA release from a single islet was estimated to reach local concentrations above 10 μ M. In conclusion, GABA secretion from islets is robust and pulsatile.

We performed experiments addressing the possibility of conventional Ca^{2+} dependent exocytosis by granules or vesicles³¹. Stimulating islets with glucose had no effect on pulsatile GABA secretion measured by biosensor cells (Figure 3d,e) or on total GABA secretion measured by HPLC (Figure 3f, Extended Data 4) (see also^{33,34}). Depolarizing islets with KCl (30 mM) or removing extracellular Ca²⁺ did not affect pulsatile GABA secretion (Figure 3g–i). Likewise, depleting extracellular calcium or simultaneously depleting intracellular Ca²⁺ sources with thapsigargin (Figure 3j), stimulating with KCl (Extended Data 4), or opening ATP-gated potassium channels with diazoxide (Extended Data 4) did not significantly affect GABA secretion from human islets as measured by HPLC. Together, these results strongly suggest that Ca²⁺ (either influx or intracellular release) is not a primary trigger for gating of GABA release from the human islet. A non-vesicular mode of GABA efflux likely depends on the cytosolic, metabolic pool of GABA^{49,50}. Inhibiting GABA biosynthesis with allylglycine (10 mM) acutely diminished beta cell GABA content and secretion (Figure 3k,m and Extended Data 5). Increasing the intracellular GABA concentration by inhibiting GABA catabolism with γ -vinyl GABA (10 μ M) increased the amount of released GABA per pulse (Figure 3l,m). Thus, the effects of manipulating GABA metabolism are surprisingly acute and suggest a high rate of GABA biosynthesis that couples GABA efflux to the cytosolic GABA pool^{34,51–53}.

VRAC and TauT transport cytosolic GABA across the plasma membrane in islet cells

In view of the high levels of non-vesicular GABA release from beta cells, we sought to determine if membrane transporters contribute to GABA efflux. We analyzed three distinct human islet single-cell RNA-seq datasets^{54–56} for expression of GABA-transporting proteins. A phylogenic tree of the neurotransmitter transporter family shows the relationship between the membrane GABA transporters (GAT1–3), the betaine-GABA transporter (BGT1), and the taurine transporter (TauT) (Figure 4a). GAT1–3 and BGT1 were not detected in beta cells (Figure 4b, Extended Data 6). However, TauT, which is also a GABA transporter^{57,58}, is highly expressed in beta cells (Figure 4b, Extended Data 6). Immunostaining confirmed expression and localization of TauT in the plasma membrane of human beta cells (Figure 4c).

We further searched for other putative GABA-transporters across the entire solute channel (SLC) gene group (395 members). Relative mRNA expression of all SLC genes in beta cells (Supplementary Table 2) revealed that the 4F2 cell-surface antigen heavy chain (4F2HC) and its heterodimer partners, LAT1 and LAT2, are highly expressed in beta cells (Figure 4b, Supplementary Table 2). We considered LAT2 as a possible GABA transporter due to its specificity for the GABA-mimetic drug gabapentin⁵⁹. Immunostaining confirmed islet-specific expression and localization of both 4F2HC and LAT2 in the plasma membrane of human islet cells (Figure 4d).

Uptake of 3H-radiolabeled GABA (3H-GABA) was measured to assess if either TauT or LAT2 mediate inward transport of GABA in human islets (Figure 4e). Competing substrates for TauT (10 mM taurine, beta-alanine) or LAT2 (10 mM leucine) were tested for inhibition of 3H-GABA uptake. Unlabeled GABA served as a positive control and glycine served as an additional control for possible GABA transport by monoamine transporters. Taurine and beta-alanine strongly inhibited uptake of 3H-GABA similarly to unlabeled GABA, while leucine and glycine were less effective (Figure 4e). Pharmacological inhibition of the membrane GABA transport family with a mixture of SNAP5114 (50 μ M), NNC05–2090 (50 μ M), and NNC711 (10 μ M) also inhibited 3H-GABA uptake. These GAT inhibitors also perturbed pulsatile GABA release measured by biosensor cells (Extended Data 4), producing an initial transient increase in extracellular GABA levels, indicating that GABA uptake is important for maintaining low interstitial levels in the islet. As we were unable to detect expression of GAT1–3 in islet cells, we propose that the GAT-blocking drugs acted on closely related TauT. Together, the data support the conclusion that TauT is the dominant mediator of GABA uptake in islets.

For GABA release, we looked for expression of known non-canonical GABA transporters, including bestrophin chloride channels (BEST1–4)⁶⁰ and VRAC⁶¹. VRAC conveys osmosensitive chloride (Cl⁻) currents and conducts efflux of GABA, taurine, and other small organic osmolytes to regulate cell volume^{61–64}. VRAC complexes are heterohexamers composed of multiple LRRC8 family subunits, LRRC8A, B, C, D or E. LRRC8A (also known as Swell1) is critical for formation of functional VRAC channels while its heteromer partners LRRC8B-E confer substrate specificity^{61,65}. VRAC subunit LRRC8D confers permeability to GABA and taurine⁶². Beta cells express VRAC^{66–68}, with channel subunits LRRC8A, LRRC8B, and LRRC8D detected by single-cell RNA-seq (Figure 4b).

To assess whether VRAC channels mediate GABA efflux, human islets were exposed to hypo-osmotic buffer and release of endogenous GABA was measured by HPLC (Figure 4f). GABA release increased by 3x or 40x upon exposure to increasingly hypotonic buffer (Figure 4f). Hypotonic induction of GABA efflux was eliminated from LRRC8A-knockout MIN6 beta cells⁶⁷ and isotonic GABA efflux was reduced by ~50% (Figure 4g–h). These results support a mechanism of cytosolic GABA release from islets via VRAC.

To assess whether VRAC in beta cells is responsible for generation of GABA release pulses we generated beta-cell specific LRRC8A knockout mice (βc -LRRC8A^{-/-}) by crossing LRRC8A floxed mice (LRRC8A^{fl/fl})^{67,69} with mice expressing Cre recombinase in insulinproducing beta cells (Ins1^{cre})⁷⁰. Loss of LRRC8A in islets isolated from βc-LRRC8A^{-/-} mice was confirmed by Western blot (Figure 4i). βc-LRRC8A^{-/-} islets do not release GABA in response to hypotonic stimulus (Figure 4j) consistent with beta cells being the dominant GABA-synthesizing cell type in the islet. In wild-type LRRC8A^{+/-} littermates GABA release was pulsatile in isotonic conditions and responded to continuous hypotonic stimulation with kinetics consistent with VRAC gating⁶²: delayed activation of GABA release of 2-3 minutes that builds to a maximum rate of release after ~8 minutes and remains active throughout the hypotonic stimulation. Biosensor cell recording of GABA release from β c-LRRC8A^{-/-} islets demonstrated both a loss of GABA pulses and loss of GABA release under hypotonic stimulus (Figure 4k-l and Extended Data 7). We next studied the effect of knocking down LRRC8A in human islets transduced with adenovirus encoding LRRC8A shRNA (Ad-mCherry-hLRRC8A-shRNA) or scrambled shRNA (AdmCherry-Scramble-shRNA) (Figure 41-n). Expression of adenovirus LRRC8A shRNA resulted in ~40% decrease in expression of LRRC8A and loss of GABA pulses (Figure 4mo). Finally, we used HPLC, a direct detection method, to validate that the kinetics of GABA released from human islets under hypotonic stimulation are consistent with previous reports of VRAC-mediated organic osmolyte release (Figure 4p)⁶². Together, these data are consistent with a critical role of the LRRC8A component of VRAC in release of GABA from the cytosol of beta cells.

Cytosolic GABA secretion synchronizes insulin secretion

GABA has been assigned many functional and regulatory roles in the islet. However, its effects on hormone secretion in human beta cells has been a matter of ongoing investigation⁷¹. We examined the autocrine effects of pulsatile GABA release on insulin secretion from human islets. Exposure of human islets at 5 mM glucose to exogenous

GABA (10μ M) reduced insulin release by ~40%, suggesting that GABA has an inhibitory effect on insulin release under normoglycemic conditions (Figure 5a). We next measured the effect of decreasing endogenous GABA release by inhibiting GABA biosynthesis with allylglycine (10μ M) (Figure 5b). Inhibition of endogenous islet GABA production increased insulin secretion during glucose stimulation (11μ M) (Figure 5b,c), consistent with an inhibitory effect of GABA on insulin secretion. Inhibition of endogenous islet GABA production did not, however, affect insulin release stimulated by depolarization with KCl (Figure 5b), indicating that allylglycine treatment did not disrupt the available pool of releasable insulin.

We next measured the effect of GABA on cytosolic Ca^{2+} responses in human islets. Human islets were loaded with Ca^{2+} indicator dye (Fluo4) and imaged by confocal microscopy during glucose stimulation (Figure 5d,e). When GABA was applied during the first phase of glucose-induced Ca^{2+} influx, cytosolic Ca^{2+} levels were quickly and dramatically reduced to near pre-stimulation levels. Islets quickly desensitized within 1–2 minutes to the inhibitory effect of applied GABA. These effects are consistent with GABA acting through GABA_A ionotropic receptors (rather than GABA_B metabotropic receptors)⁷². GABA applied to islets in resting glucose conditions (3 mM) had no effect on beta cell Ca^{2+} calcium responses.

To test the effects of GABA on coordinated pulsatile insulin secretion, we measured serotonin release as a surrogate for insulin secretion^{73–77} using CHO cells expressing the serotonin receptor 5-HT2C^{78,79}. As we and others have reported^{77,79–81}, serotonin/insulin secretion from human islets was pulsatile with regular periods ranging from 4 to 10 minutes (Figure 5f). When we decreased endogenous GABA levels and release from cells by inhibiting GABA biosynthesis with allylglycine (10 mM), basal serotonin/insulin secretion increased and failed to display regular secretory pulses (Figure 5f,h). Decreasing catabolism of GABA and increasing endogenous GABA levels by adding γ -vinyl-GABA (10 μ M), inhibited pulsatile serotonin/insulin secretion (Figures 5g,h). Similarly, blocking GABAA receptor signaling with the inhibitor SR 95531 (10 μ M) disrupted the periodicity of serotonin pulses (Figure 5i). These results indicate that GABA production and paracrine GABAergic signaling within the islet impacts periodic insulin secretion.

By expressing the lumenal protein of large dense core vesicles neuropeptide Y (NPY) fused to the pH-dependent green fluorescent protein pHluorin in beta cells, we visualized exocytotic events in real time^{82,83} and observed periodic exocytotic bursts occurring simultaneously throughout the islet with periods of ~ 5 minutes. In the presence of the GABA_A receptor agonist muscimol (100 μ M) these exocytotic events became smaller and lost synchronicity (Figures 5j,k).

Thus, using four different methods, we provide evidence that endogenously produced GABA released from a cytosolic beta cell pool decreases insulin release while stabilizing the periodicity and glucose responsiveness of insulin secretion.

Cytosolic GABA secretion is interrupted in human islets from type 2 diabetic donors

As shown in Figure 1, immunostaining of human islets from type 2 diabetic donors revealed depletion of GABA pools in beta cells in spite of robust expression of the biosynthesizing

enzyme GAD65. We examined multiple preparations from human type 2 diabetic donors for GABA release with biosensor cells (n 3 islets from each of the 5 donors). Consistent with the immunostaining results, we could not detect pulsatile GABA secretion from human islets from type 2 diabetic donors (Figure 6a,b). Type 2 diabetic islets did respond to KCl depolarization as evidenced by strong increase in cytosolic Ca²⁺, indicating they were alive and retained membrane potentials (Figure 6c). By contrast, pulsatile GABA secretion was robust in islets from non-diabetic donors (Figure 6a, see also Figure 3). Pulsatile GABA secretion from islets of type 2 diabetic patients could be rescued by inhibiting GABA catabolism with γ -vinyl GABA (10 μ M) (Figure 6d–e). The periodicity of insulin release, measured by serotonin co-release detected via biosensor cells, was deranged in type 2 diabetic islets (Figure 6f,g). This effect was similar to that elicited by blocking GABA signaling in non-diabetic islets by inhibiting GABA_A receptors with SR 95531 (Figure 5i). Insulin secretion periodicity in type 2 diabetic islets became more regular upon treatment with γ -vinyl GABA (Figure 6g).

We previously reported the accumulation of GAD65 in Golgi membranes of beta cells undergoing ER stress through perturbations of the palmitoylation cycle that controls targeting of the enzyme to peripheral vesicles⁸⁴. Similar accumulation and defect in membrane compartment distribution was detected in individuals experiencing early as well as late phases of GAD65 autoimmunity and development of type 1 diabetes⁸⁴. We examined the intracellular distribution of GAD65 by confocal microscopy of immunostained sections of human pancreas. GAD65 was detected in both the cytoplasm and Golgi compartment in beta cells in sections from non-diabetic donors (Figure 6h), but showed increased localization to the Golgi compartment in beta cells of type 2 diabetic patients (Figure 6i,j).

DISCUSSION

Our study provides evidence for a novel mechanism of GABA release in human beta cells. As a result of enzymatic synthesis of GABA from glutamic acid by GAD65, GABA is present at high levels in the cytoplasm of beta cells. It is from this cytosolic pool that GABA is released, because only ~1% of the beta cells show evidence of VGAT expression and the consequent accumulation of GABA in secretory vesicles. Our study shows that GABA in the islet behaves as an organic osmolyte. Indeed, in the islet, GABA shares transport properties with the canonical organic osmolyte taurine, namely efflux *via* VRAC and uptake *via* the taurine transporter. We further found that islet GABA levels are greatly reduced under diabetic conditions. Given the paracrine, islet-trophic, and immunosuppressive roles of GABA, the loss of islet GABA content in both type 1 and type 2 diabetes may contribute to beta cell loss and dysfunction.

Several mechanisms can be suggested to cause the low levels of islet GABA in human diabetes. First, while the GAD67 isoform binds the co-enzyme 5' pyridoxal phosphate (PLP) firmly and is a constitutively active holoenzyme, GAD65 oscillates between an active holoenzyme and an inactive apo-enzyme^{85,86}. Thus, it is possible that GAD65 is mainly present as an inactive apo-enzyme in beta cells under diabetic conditions. Second, GAD65 may be rendered inactive in beta cells by expression or formation of an inhibitor. Third,

GABA metabolism, rather than GAD65 expression, may be a dominant factor controlling islet GABA content in beta cells in diabetic conditions.

GAD65 anchors to the cytosolic face of intracellular membranes²⁵. The shift in localization of the enzyme to perinuclear ER/Golgi membranes observed in beta cells in human diabetes would not be expected to prevent release of the product GABA into the cytosol unless the enzyme in Golgi membranes represents the inactive GAD65 apoenzyme or is associated with an inhibitor. The consequent decrease in synthesis of cytosolic GABA would interrupt GABA transport and prevent pulsatile GABA secretion. This possibility would be consistent with our observations in human alpha cells, which are devoid of GABA, yet express GAD65 that appears to localize exclusively to ER/Golgi membranes.

Expression of the synaptic vesicle markers VGAT and synaptophysin in islets^{87,88} has contributed to the concept that GABA is secreted from beta cells *via* synaptic-like microvesicles²⁸. Here, using an approach that allows for high resolution subcellular localization studies, we show that almost all beta cells (> 99%) lack synaptic-like vesicles or granules containing VGAT together with GAD65 and/or GABA. A small subset of beta cells (< 1%) exhibit all features of vesicular GABA. The nature of the small subpopulation of VGAT-positive beta cells is currently unknown but may intersect with markers of beta cell subtype and/or maturation state. It is possible that its size is variable and subject to presently unknown factors which could affect its contribution to the vesicular release of GABA described earlier^{26,31,89}. However, it appears that in the vast majority of beta cells, GABA is released from the cytosolic pool *via* plasma membrane channels.

Pertinent to our findings, Rorsman and Pipeleers have previously reported a high rate of basal (non-quantal) GABA release that was unregulated by glucose or pharmacological regulators of insulin secretion^{31,34,90}. Due to this unregulated GABA release, it was concluded that the beta cell must be equipped with a second pathway for release of GABA that is non-vesicular, the details of which remained to be elucidated^{31,90}. Here, we have identified VRAC to be a pathway for GABA release that is consistent with the findings by Rorsman and Pipeleers.

While glucose-inducible VRAC Cl⁻ currents have been observed in beta cells^{67,68}, no effect of glucose has been reported for LRRC8A-dependent organic osmolyte efflux. This may be because LRRC8A-dependent Cl⁻ and organic osmolyte efflux are not required to follow the same behavior. For example, organic osmolyte and Cl⁻ currents can occur through different isoforms of the VRAC channel. LRRC8A/D is the dominant channel for GABA release and LRRC8A/B/C/E channels exhibit low GABA conductance but high Cl⁻ conductance^{62,65}. It remains to be determined if multiple isoforms of VRAC exist simultaneously in beta cells or if the same VRAC channel can differentially gate Cl⁻ and organic osmolytes depending on the activating conditions. Another explanation for why we do not observe glucose-mediated effects on GABA release may be that the degree of swelling required to activate the GABA VRAC channel is greater than obtained by glucose-stimulation. The Jentsch group reported that a very high, non-physiological glucose stimulation (25 mM = 450 mg/dl) induced comparatively moderate beta cell swelling (only 1/4 of the volume differential of hypotonic stimulation) and no clear subsequent regulatory volume decrease⁶⁸. The Sah group obtained

similar results, where they showed that 16.7 mM glucose induces only a minor beta cell swelling of 6.8% in murine beta cells with a sluggish associated volume response, and no clear trend of glucose on human beta cell swelling⁶⁷. If regulatory volume decrease is required for organic osmolyte efflux via VRAC, then this lack of a regulatory volume decrease upon glucose stimulation is consistent with our observation that glucose does not induce GABA release.

Our findings indicate that endogenously released GABA has two major effects on beta cells: (A) it reduces insulin secretion and (B) it helps stabilize the periodicity of insulin pulses. That GABA has inhibitory effects on beta cells was further supported by experiments in which GABA and the GABAA receptor agonist muscimol were added exogenously and is consistent with findings by Birnir and colleagues⁷¹. Similar to the role somatostatin is proposed to play in the islet^{91,92}, GABA may serve to change the gain of insulin secretion and thus prevent its wasteful release. The pulsatile pattern of GABA efflux and its impact on the periodicity of insulin secretion suggest an additional role for GABA in timing or pacing of oscillatory islet activities. As shown by our results, restoring GABA signaling in type 2 diabetic islets improves the synchronicity of insulin secretory pulses but diminishes their magnitude. Conversely, the loss of GABA in diabetic states is likely to produce increases in the excitability of beta cells that will help increase insulin secretion, but at the price of losing periodicity. While continuous pulsatile GABA release may contribute to the economy and periodicity of insulin secretion under normal conditions, it remains to be determined if the dramatic reduction in GABA levels is a mechanism that helps the beta cell increase insulin secretion and hence cope with the increased demand in diabetic states.

Our findings that GABA inhibits insulin secretion and beta cell Ca²⁺ responses differ from those showing that GABA depolarizes beta cells and increases insulin secretion³¹. The discrepancy can be explained by different experimental conditions (e.g. dynamic hormone secretion measurements *versus* static incubation, different basal glucose concentrations). That GABA has been reported to depolarize beta cells to $\sim -50 \text{ mV}^{31}$, however, indicates that GABA will clamp the membrane potential below the threshold for the opening of P/Q type Ca²⁺ channels (above -20 mV), the channels that are responsible for insulin granule exocytosis⁹³. Our findings are further in line with results showing that GABA stimulates delta cells³¹, which leads to secretion of the potent inhibitory hormone somatostatin. Importantly, beta cells also express inhibitory metabotropic GABA_B receptors whose activation opens hyperpolarizating K⁺ channels or inhibit adenylyl cyclases⁹⁴. Therefore, there is substantial evidence supporting an inhibitory role for GABA in human beta cells.

It is likely that many physiological processes within the islet, including local actions of secreted GABA and of other paracrine signals^{31,43,44}, shapes islet cell excitability, as described for neurons in the central nervous system^{36,95}. In addition to the effects on insulin secretion reported here, non-vesicular GABA secretion also affects the activities of the glucagon-secreting alpha cell and the somatostatin-secreting delta cell^{27,96}. Because GABA stimulates delta cells to secrete somatostatin and because somatostatin strongly inhibits insulin secretion, a pulse of GABA may both inhibit beta cells directly and indirectly via delta cells. Recent findings by the Sah and Jentsch groups demonstrate that knockout of LRRC8A, the GABA releasing pathway we report here, impairs glucose-responsive insulin

secretion in contrast to our finding that GABA is inhibitory to beta cells^{67,68}. A possible explanation for this discrepancy is that insulin secretion is an event requiring integration of multiple signals. Sah and Jentsch both showed that loss of LRRC8A also affects beta cell membrane potential and delays or impairs beta cell Ca²⁺ responses. As GABA appears to only be inhibitory to beta cells during glucose-responsive Ca²⁺ fluxes, any loss of inhibition from impaired GABA release may be overwhelmed by the stronger inhibition imposed by LRRC8A knockout. Thus, the functional effect of a loss of GABA is observable by blocking GABA biosynthesis, as we have shown, but not by preventing GABA release through VRAC knockout.

Intracellular GABA levels and cytosolic GABA release are dramatically decreased in type 1 (Figure 1) and type 2 diabetes (Figures 1, 6), indicating that islets lose the paracrine, trophic, and immunomodulatory influence of GABA in the diabetic state. It is conceivable that this loss of GABA leaves islets vulnerable to destructive inflammation. We propose that a periodic pattern of cytosolic GABA release independent of glucose concentration impacts the magnitude and periodicity of insulin secretion. Interrupting GABA secretion impairs coordination of hormone secretion. Irregular insulin secretion from the islet may exacerbate insulin resistance⁴⁸. Given its inhibitory effect on alpha cells, defective GABA signaling⁹⁷ and decreased GABA secretion could also explain why glucagon secretion is increased in type 2 diabetes, causing further elevation of hyperglycemia. Thus, loss of GABA signaling in the islet may contribute to the pathogenesis of type 1 and 2 diabetes. Because restoring GABA signaling can be proposed as an intervention point to promote islet function, our study has implications for a novel pharmaceutical strategy for the treatment of diabetes. In rodent islets electrical coupling via gap junctions and purinergic paracrine signaling have been suggested to coordinate rhythmic insulin secretion⁴⁸. Here we show, however, that in the human islet GABA is a potential pacemaker candidate because (1) it is released independently of glucose concentration in pulses with a frequency in the range of those of pulsatile in vivo insulin secretion; (2) it is a diffusible factor acting on GABAA receptors whose activation inhibits beta cell activity; and (3) its production and release can regulate the periodicity of insulin secretion.

METHODS

Human pancreas tissues

Human pancreatic sections from tissue donors of both genders were obtained via the Network for Pancreatic Organ Donors with Diabetes (nPOD) tissue bank, University of Florida, Gainesville, FL, USA. Human pancreata were harvested from cadaveric organ donors by certified organ procurement organizations partnering with nPOD in accordance with organ donation laws and regulations and classified as "Non-Human Subjects" by the University of Florida Institutional Review Board (IRB No. 392–2008) waiving the need for consent^{98,99}. nPOD tissues specifically utilized for this project were approved as non-human by the University of Florida Institutional Review Board (IRB No. 201701113).

Human pancreatic islets were obtained from deceased non diabetic donors and from donors with type 2 diabetes from the Human Islet Cell Processing Facility at the Diabetes Research Institute at the University of Miami Miller School of Medicine, from the NIDDK-funded

Integrated Islet Distribution Program (IIDP) at City of Hope, from the European Consortium on Islet Transplantation (ECIT) Islets for Basic Research Program, and from Prodo Laboratories, Inc. Human pancreatic islets from deceased donors with type 1 diabetes were isolated by the nPOD Islet Isolation Program (IIP). Human islets received from the University Hospital of Geneva and San Raffaele Scientific Institute, Milan through the ECIT islets for basic research program were approved by the Institutional Review Board of the University Hospital of Geneva (CER No. 05–028) and by the Ethics Committee of the San Raffaele Scientific Institute of Milan (IPF002–2014). The University of Geneva and the San Raffaele Institute Ethics Committees waived the need for consent from the donors because islets were used for experimental research only when not suitable for clinical purposes and would otherwise have been destined for destruction. In such cases obtaining informed consent is not mandatory in Switzerland and Italy. Cadaveric human islets for research were approved as non-human by the University of Florida Institutional Review Board (IRB No. 201702860).

Human islets were cultured at 24°C in 10 cm non-adherent cell culture dishes (500 islets/ dish) in CMRL medium with 2% glutamine, 10% FBS, 10 mM HEPES and 1% Penicillin/ Streptomycin.

Rat pancreatic islets

All experimental protocols using rat islets were approved by the University of Florida and EPFL Animal Care and Use Committees. Rat islets were isolated from pancreases of male and female P5 Sprague Dawley rats (Charles River) in accordance with published methods¹. Rat pups were sacrificed by decapitation and the whole pancreas was removed and digested in 0.15 mg/ml Liberase TL (Roche #05401020001) in HBSS (Hank's buffered salt solution, Gibco #24020091) + 20 mM HEPES for 7 minutes with strong manual agitation. The Liberase enzyme was stopped with addition of HBSS + 20 mM HEPES + 0.5% fetal bovine serum (FBS). Digested pancreas tissue was washed four times with HBSS + 20 mM HEPES + 0.5% FBS at 4°C, resuspended in Histopaque-1119 (Sigma #11191), overlain with HBSS + 20 mM HEPES + 0.5% FBS (room temperature), and centrifuged for 20 min at 300 x g at 20°C. Islets were collected from the interface between Histopaque and HBSS phases and washed 3x with HBSS + 20 mM HEPES + 0.5% NBCS. Islets were then hand-cleaned with a 200 µl pipette and cultured in 10 cm non-adherence petri dishes at 37°C 5% CO₂, 500 islets per dish, 9 ml RPMI 1640 medium with GlutaMAX (Gibco #61870010), 10% FBS, 1% Penicillin/Streptomycin.

LRRC8A knockout mouse islets

All experimental protocols using transgenic mice were approved by the University of Florida Animal Care and Use Committee. Experimental protocols using transgenic mouse islets were approved by the University of Florida Animal Care and Use Committee. *LRRC8A*-floxed (*LRRC8A*^{fl/fl}) mice on the C57BL/6NCrl background were generated in and provided by Rajan Sah's labs at the University of Iowa and Washington University in St. Louis^{67,69}. *LRRC8A*^{fl/fl} mice were crossed with mice expressing Cre recombinase in beta cells (Ins1^{cre})⁷⁰ purchased from The Jackson Laboratory (B6(Cg)-*Ins1^{tm1.1(cre)Thor/JI*; Stock #026801) to generate beta cell specific LRRC8A-knockout (βc-LRRC8A^{-/-}) mice. Mouse}

genotypes were confirmed by PCR using published primers^{67,69,70} and loss of LRRC8A in islets was confirmed by Western blot. βc -*LRRC8A*^{-/-} mice (8–14 weeks old) were sacrificed by cervical dislocation under deep isoflurane anesthesia according to the approved procedures. Heterozygous *LRRC8A*^{+/-} Cre-expressing littermates served as wild-type controls. Equal numbers of male and female mice were used. The pancreas was perfused via the common bile duct with 2–3 ml HBSS containing Liberase TL (0.15 mg/ml), removed, and digested at 37 °C for an initial 12 minutes, disrupted by pipetting up and down several times with a 10 mL pipette, and digested for a final 4 minutes. The enzyme digestion was quenched with HBSS + 10% FBS. Islets were purified from digested tissue and cultured using the same methods as described for rat islets.

Islet cell monolayer culture

Within one week of isolation, rat or human islets were hand-picked from suspension cultures, collected in a 15 ml tube and washed twice in PBS without Ca^{2+} and Mg^{2+} . Islets were dissociated into a suspension of single islet cells by continuous gentle pipetting in 0.3 ml 0.05% trypsin-EDTA per 500 islets for 3 minutes at 37 °C. Trypsin digestion was halted by addition of islet monolayer medium (Minimum essential medium (MEM) with GlutaMAX 11 mM glucose, 5% FBS, 1 mM sodium pyruvate, 10 mM HEPES and 1x B-27 Supplement) to a total volume of 15 ml, followed by pelleting of islet cells by centrifugation for 5 min at 1400 rpm ($350 \times g$) and resuspension islet monolayer medium. Islet cells were seeded on round 12 mm diameter and 0.17 mm thickness borosilicate glass coverslips (Electron Microscopy Sciences) coated with purified laminin (Gibco) or purified collagen IV (Sigma Aldrich) at 50 µg/ml in HBSS with Ca^{2+}/Mg^{2+} for 1 hour at 37 °C. Cells were seeded at approximately 35,000 cells/cm². Islet cells required 3–4 days of culture to adhere and spread on surfaces before further experimentation. A detailed description and validation of monolayer cultures of primary human and rat islets cells is available¹⁰⁰.

Hippocampal neuron cultures

Primary rat hippocampal neurons were prepared from P2-P3 Sprague Dawley rats of both sexes, as described by Codazzi, et al¹⁰¹. Neurons were seeded on round poly-L-ornithine-coated glass or Thermanox coverslips (Nunc), at 100,000 cells per coverslip in a 24-well plate and in 1 ml of neuronal medium: Minimum essential medium (MEM) with GlutaMAX (Gibco), 11 mM glucose, 5% FBS, 1 mM sodium pyruvate, 10 mM HEPES and 1x B-27 Supplement53 (Gibco). One day after isolation, 3 μ M of the chemotherapeutic agent ARA-C (Sigma-Aldrich) was added to the culture medium to eliminate astrocytes and obtain a neuronal culture of high purity (>90% neurons).

Immunofluorescence staining

Human pancreas sections obtained from nPOD were deparaffinized followed by acidic-pH heat-mediated antigen retrieval according to the nPOD standard operating procedure for immunopathology. Monolayers of pancreatic islet cells were fixed with 4% EM-grade PFA (Electron Microscopy Sciences) at room temperature for 20 minute. Samples were blocked and permeabilized in PBS + 0.3% Triton X-100 with 10% goat or donkey serum. Primary antibodies were incubated overnight in PBS + 0.3% Triton X-100 with 1% goat or donkey serum at 4°C. Alexa Fluor 405, 488, 568, and 647 conjugated secondary antibodies (Thermo

Fisher) were incubated at 1:200 dilution in PBS + 0.3% Triton X-100 for 30 minutes at room temperature. Coverslips were mounted with ProLong Gold Antifade Reagent with or without DAPI (Thermo Fisher).

Immunostaining for GABA and taurine was validated by competitive inhibition of primary antibody binding by addition of soluble GABA or taurine to the antibody incubation buffer (Extended Data 6). The anti-GABA and anti-taurine antibodies showed minimal non-specific binding towards non-GABA or non-taurine amino acids. Furthermore, GABA immunostaining was eliminated by inhibition of the GABA-synthesizing enzyme, GAD65, with allylglycine (Extended Data 5).

Microscopy

Confocal images (pinhole = airy 1) of randomly selected islets (2-3 islets per section) were acquired on a confocal laser-scanning microscope (Zeiss LSM700, Zeiss LSM710, Leica SP5, and Leica SP8) with 20x/0.8 NA Plan-Apochromat air, 40x/1.30 and 63x/1.40 NA Plan-Apochromat oil-immersion objectives at 1024×1024 pixel resolution. Images were processed and quantified in ImageJ. Manders' coefficient colocalization analyses were performed using the JACoP (Just Another Colocalization Plugin) plugin for ImageJ¹⁰². To determine the subcellular localization of GAD65, the intensity of GAD65 immunostaining in the Golgi compartment of beta cells in pancreata from healthy donors was compared to that of pancreata from T2D donors. Using ImageJ software, we measured GAD65 staining intensity in the whole cells as well as in the Golgi apparatus by selecting a region of interest based on the Golgi staining signal. GAD65 immunostaining was always more intense in the Golgi apparatus than in the cytosol. To allow comparisons between tissue sections and specimens, we expressed the mean GAD65 staining intensity in the Golgi compartment relative to the intensity of GAD65 staining in the whole beta cell [GAD65 = (mean GAD65 intensity in Golgi - mean GAD65 intensity in whole cell) / mean GAD65 intensity in whole cell].

Gene expression analysis

Raw molecular counts per gene and cell were directly obtained from GEO (accession numbers GSE84133, GSE81076, and GSE83139), and further normalized using the R package scran¹⁰³. Results were qualitatively similar across the three datasets. Log(norm_values+1) corresponding to GSE81076 are displayed in Figure 4 and GSE84133 and GSE83139 in Extended Data 6.

Detection of GABA via biosensor cells

We adapted real time measurements of GABA secretion from Dvoryanchikov et al¹⁰⁴. GABA biosensor cells were obtained from Novartis Institutes for BioMedical Research in Switzerland. GABA biosensor cells consisted of Chinese hamster ovary (CHO) cells stably expressing heteromeric GABA_B (GABA_B R1b and GABA_B R2) receptors and the G-protein a subunit, Gaqo5 modified to couple to increases in $[Ca^{2+}]_i$ via the InsP₃ signaling cascade (Extended Datas 3,7). GABA biosensor cells reliably responded to low concentrations of GABA (threshold \approx 100 nM), making them highly sensitive GABA detectors. GABA biosensor cell responses were measured using $[Ca^{2+}]_i$ imaging. We loaded GABA

biosensors with the $[Ca^{2+}]_i$ indicator Fura-2 and plated them on poly-d-lysine coated cover slips in a perfusion chamber. Individual human islets were placed on top of this layer of biosensor cells. Fluid perfusion was performed with a gravity-driven Warner Instruments VC-8 8-channel perfusion system set to 0.5-1 ml/min and connected to a Warner Instruments RC-20 closed bath small volume imaging chamber to ensure linear solution flow and fast exchange. GABA secretion was examined in biosensors cells located immediately downstream of the islet in recordings lasting at least 20 minutes to be able to detect rhythmic behavior. GABA secretion was examined for pulses simply by inspecting recordings for robust increases in [Ca²⁺]; in biosensor cells. Because the pulse amplitudes were large, no deconvolution or other processing of the raw $[Ca^{2+}]_i$ traces was necessary. We calculated the periods between pulses by measuring the time between the initial rises in $[Ca^{2+}]_i$ in at least three sequential pulses per biosensor cell. The regularity of the pulses was quantified by using the deviation of this interpulse interval. Changes in the amount of secreted GABA were quantified by measuring the area under the curve of the $[Ca^{2+}]_i$ responses in the biosensor cells during defined time intervals. These analyses were only performed within the same experiment because quantitative comparisons between experiments would have required calibration with known concentrations of GABA. Only recordings with at least three responsive biosensor cells were included in the analyses. Secretion was considered coordinated and pulsatile if the responses in the biosensor cells were synchronized and showed regular periods. We expressed these data as average traces of the $[Ca^{2+}]_i$ responses of the biosensor cells.

We established that biosensor cells responded only to GABA and not to other substances including taurine. Of all the tested substances, only GABA activated the biosensor. Crucially, the antagonist CGP (10 µM) completely blocked GABA_B receptors on the GABA biosensors and eliminated responses generated from islets. Stimuli or pharmacological agents (e.g. antagonists, transporter blockers) used in this study did not themselves either elicit biosensor responses or alter the ability of biosensors to respond to GABA. We conducted these controls by [Ca²⁺]_i imaging of biosensor cells plated at low density and in the absence of islets. When examining pulsatile secretion it was important to establish that biosensor cells themselves did not display periodic behavior in the absence of islets or in the continuous presence of GABA. Biosensor cells for acetylcholine, which are also CHO cells, did not show oscillatory responses in the presence of islets⁴⁴, indicating that oscillatory signals are not an intrinsic property of these cells but stem from the islet's secretory behavior. The effects of manipulation were compared to controls recorded in the same experimental session or using islets from the same human islet preparation to compensate for the variability in the quality of islets. To ensure that islets were healthy, we simultaneously monitored [Ca²⁺]; responses in islets and biosensor cells. KCl depolarization induced responses in islets but not in biosensor cells, indicating that islet cells were viable. False negative results were ruled out by confirming that biosensor cells remained fully responsive to GABA at the end of the recording session.

Detection of insulin via biosensor cells

To detect insulin release we used an approach in which serotonin is used as a surrogate for insulin. Serotonin is present in insulin granules and is released with insulin^{73–77,81}.

Biosensor cells for serotonin were CHO cells expressing the serotonin receptor 5-HT_{2C} and are further described and characterized in previous publications^{78,79}.

Determination of cytosolic Ca²⁺ concentration

Imaging of cytoplasmic $[Ca^{2+}]$ ($[Ca^{2+}]_i$) was performed in accordance with published descriptions¹⁰⁵. Islets, dispersed islet cells, or biosensor cells were incubated in Fura-2 AM (2 μ M; 1 hour) and placed in a closed small-volume imaging chamber (Warner Instruments, Hamden, CT). Stimuli were applied with the bathing solution. Cells loaded with Fura-2 were alternatively excited at 340 and 380 nm light and fluorescence was recorded on two different microscope setups. At the University of Miami, we used a monochromator light source (Cairn Reseach Optoscan Monochromator, Cairn Research Ltd, Faversham, UK). Images were acquired with a Hamamatsu camera (Hamamatsu Corp, Japan) attached to a Zeiss Axiovert 200 microscope (Carl Zeiss, Jena, Germany). Changes in the 340/380 fluorescence emission ratio were analyzed over time in individual cells using MetaFluor imaging software. At the University of Florida, we used a pE-340fura LED illumination system (CoolLED Ltd., Andover, UK) and a Hamamatsu ORCA-Flash 4.0 LT+ camera attached to a Zeiss Axio Observer Z1 microscope. Change in the 340/380 fluorescence emission ratio were analyzed over time in individual cells using MetaFluor imaging software.

Insulin secretion during perifusion

A high-capacity, automated perifusion system was used to dynamically measure insulin secretion from pancreatic islets (Biorep Perifusion V2.0.0, Miami, FL). A low pulsatility peristaltic pump pushed KRBH solution at a perifusion rate of 100 μ L/min through a column containing 100 pancreatic islets immobilized in Bio-Gel P-4 Gel (BioRad, Hercules, CA). Except otherwise stated, glucose concentration was adjusted to 3 mM for all experiments. Stimuli were applied with the perifusion buffer. The perifusate was collected in an automatic fraction collector designed for a 96 well plate format. The columns containing the islets and the perifusion solutions were kept at 37°C, and the perifusate in the collecting plate was kept at < 4°C. Perifusates were collected every minute. Insulin release in the perifusate was determined with the human or mouse Mercodia Insulin ELISA kit following manufacturer's instructions.

KRBH Buffer preparation

Kreb's ringer bicarbonate HEPES (KRBH) buffer (115 mM NaCl, 4.7 mM KCl, 2.5 mM CaCl₂, 1.2 mM KH₂PO₄, 1.2 mM MgSO₄, 25 mM NaHCO₃, 25 mM HEPES, 0.2% BSA, 3 mM glucose) was prepared containing final concentrations of solutes according to Supplementary Table 3. Theoretical osmolarity was calculated according to the following expression¹⁰⁶:

$$osmolarity = \sum_{i} \varphi_i \, n_i \, C_i$$
 Equation 1.

Where: φ is the osmotic coefficient, *n* is the number of particles (e.g. ions) into which a molecule dissociates, *C* is the molar concentration of the solute, and *i* is the identity of a particular solute. Osmolarities of solutions were verified using an osmometer and found to

be in good agreement with theoretical calculations. Isotonic KRBH had an osmolarity of 294 mOsmol/L.

Detection of GABA by HPLC with electrochemical detector

Fractions collected from islet perifusion in the BioRep islet perifusion device or supernatants from static incubations in KRBH buffer were analyzed for GABA content using an EICOM HTEC-500 HPLC-ECD with autosampler, online automated OPA-derivatization, and Eicompak FA-3ODS separation column. This automated detection technique is linearly sensitive for GABA from the nano-molar to milli-molar range¹⁰⁷. Insulin content in the same sample fraction was determined by ELISA kit (Mercodia). Data shown in Figure 3f analyzed GABA content from perifusion fractions while data shown in Figures 3j, 4f,h,j, and Extended Data 4a,b,c are from static incubations. Each sample (for all cases) contained approximately 100 IEQ. Perifusion flow rate was 100 μ L/min. Static incubations were performed in 100 μ L KRBH 3 mM glucose for 30 minutes.

LRRC8A knockout MIN6 cells

Wild-type and *LRRC8A* (also known as *Swell1*) knockout MIN6 beta cells generated by CRISPR/Cas9 technology⁶⁷ were provided by Rajan Sah's lab at the University of Iowa and Washington University in St. Louis. Confirmation of *LRRC8A* gene disruption by PCR, *LRRC8A* protein deletion, and ablation of LRRC8A-mediated current in these cells was published by the Sah group⁶⁷. MIN6 cells cultured in DMEM with 15% FBS and 1% penicillin streptomycin were transfected with human GAD65-GFP plasmid³⁹ using Lipofectamine 2000.

Adenovirus

Human adenoviruses type 5 with hLRRC8A-shRNA (Ad5-mCherry-U6-hLRRC8A-shRNA) and a scrambled non-targeting control (Ad5-U6-scramble-mCherry) were obtained from Vector Biolabs. Adenovirus was added to human islets in culture (final concentration of 5×10^7 PFU/ml) and incubated for 24 h. The islets were then washed with PBS three times and cultured for 1–2 days before performing further experiments. Transduction efficiency was assessed by fluorescence microscopy.

Western blotting

Cell lysates were prepared by extraction of whole islets or MIN6 cells in RIPA buffer (Sigma). The BCA protein assay kit (Thermo Fisher Scientific) was used to measure the protein concentration of cell extracts. Gel electrophoresis was performed with the NuPAGE system (Life Technologies) with transfer onto polyvinylidene fluoride membranes with the iBlot 2.0 (Life Technologies) device. Membranes were blocked with 5% nonfat milk in tris buffered saline, incubated in primary antibody overnight at 4°C, and detected with secondary antibody (LI-COR Biosciences). Blots were imaged on the LI-COR Odyssey CLx scanner.

Real-time recording of exocytosis

To image exocytosis, an adenovirus was engineered, encoding for an endogenous protein of large dense core vesicles neuropeptide Y (NPY) fused to pHluorin, a pH-dependent green fluorescent protein¹⁰⁸. Human islets infected with this virus were cultured short term (1 week) to permit exogenous adenoviral protein expression while retaining islet cell function. The NPY-pHluorin fusion protein was correctly localized to granules, and the pH-dependent fluorescence of pHluorin was retained. The NPY-pHluorin fusion protein exploits the granule luminal pH changes that occur during exocytosis to visualize exocytotic events of live islet cells in real time with high spatial resolution in three dimensions⁸².

Statistical Analysis

All measurements were taken from distinct samples. Means among three or more groups were compared by analysis of variance (ANOVA) in GraphPad Prism 8 software. If deemed significant, Tukey's post-hoc pairwise comparisons were performed. Means between two groups were compared by two-tailed Student's *t*-test. Variances between two groups were compared by F test. A confidence level of 95% was considered significant. The statistical test used, exact P-values, and definition of n, are all indicated in the individual figure legends. All error bars in the figures display the mean \pm s.e.m.

Reagents

The following antibodies were utilized for immunofluorescence staining:

gp-a-insulin (Linco #4011–01), gp-a-insulin (Dako #A0564), ck-a-insulin (Abcam #ab14042), sh-a-glucagon (Abcam#ab36232), sh-a-somatostatin (Abcam #ab35425), rb-a-pancreatic polypeptide (Abcam #ab113694), rb-a-GABA (Sigma-Aldrich #A2052), ms-a-GAD65 C-term¹⁰⁹ (in-house RRID:AB_528264), ms-a-GAD65 N-term¹¹⁰ (Christiane Hampe Lab), rb-a-GAD65 (Synapic Systems #198102), gp-a-GAD65 (Synaptic Systems #198104), rb-a-TauT (Sigma-Aldrich #HPA015028), rb-a-VGAT Oyster 650-labeled (Synaptic Systems #131103C5), rb-a-synaptophysin (Abcam #ab14692), rb-a-SV2C (Synaptic Systems #119202), rb-a-4F2HC (Santa Cruz Biotech. #sc9160), ms-a-LAT2 (OriGene #UM500058), rb-a-Bestrophin 1 (Abcam #ab14928), rb-a-GAT1 (Synaptic Systems #274102), rb-a-GAT2 (Abcam #ab2896), rb-a-GAT3 (Synaptic Systems #274303), rb-a-GCP60 (Novus Biologicals # NBP1–83379), rb-a-taurine (Sigma-Aldrich #AB5022), Alexa Fluor conjugated secondary antibodies (Thermo Fisher).

The following antibodies were utilized for Western blotting:

rb-α-LRRC8A (Cell Signaling #24979), ms-α-beta actin (Sigma-Aldrich #A1978), IRDye conjugated secondary antibodies (LI-COR, Inc).

gp = guinea pig, ck = chicken, ms = mouse, rb = rabbit, sh = sheep

The following chemicals were utilized:

CGP 55845 hydrochloride (Tocris Bioscience #1248/10), γ -vinyl GABA (Tocris #0808), Lallylglycine (Sigma-Aldrich #A7762), SR 95531 hydrobromide (Tocris #1262), Niflumic

acid (Tocris #4112/50), CFTRinh 172 (Tocris #3430/10), NNC711 (Tocris #1779/10), SNAP 5114 (Tocris #1561/10), NNC05–2090 hydrochloride (Tocris #2747/10), Tetrabenazine (Tocris #2175), Diazoxide (Tocris # 0964), GABA (Sigma-Aldrich #A5835), Serotonin (Sigma-Aldrich #14927), DIDS (Sigma-Aldrich #D3514), 3H-GABA (Perkin-Elmer #NET191250UC), Muscimol (Tocris #0289), Liberase TL (Roche #05401020001), Histopaque-1119 (Sigma-Aldrich #1191)

Reporting Summary

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

Data availability

The unique biological materials used in the manuscript are available from the corresponding authors upon reasonable request with the exception of those materials that the authors obtained via a materials transfer agreement (MTA) that prohibits transfer to third parties; these include the GABA biosensor cells (obtainable from Dr. Klemens Kaupmann, Novartis Institute for BioMedical Research, Basal, Switzerland), LRRC8A^{-/-} MIN6 cells and LRRC8A^{fl/fl} mice (obtainable from Dr. Rajan Sah, Washington University in St. Louis, U.S.A.), and NPY- pHluorin (obtainable from Dr. Herb Gaisano, University of Toronto, Canada). Other requests for materials should be addressed to corresponding authors Drs. Steinunn Baekkeskov, Alejandro Caicedo or Edward Phelps.

The data that support the findings of this study are available from the corresponding authors upon reasonable request.

Extended Data



Extended Data Fig. 1. Delta cells in human islets contain GABA and express GAD65; taurine content is preserved in diabetic islets

a-b. Islets in a non-diabetic human pancreas immunostained for GABA and somatostatin (a); or GAD65 and somatostatin (b). GABA and GAD65 are present in somatostatin producing human delta cells. Scale bar 50 μ m. Right panels show higher magnification views of the boxed region showing channels for: (a) (1) GABA only, (2) somatostatin only, and (3) GABA and somatostatin; (b) (1) GAD65 only, (2) somatostatin only, and (3) GAD65 and somatostatin. Images are representative of data plotted in Figures 1b and 1j. Scale bar 20 μ m.

c. Quantification of taurine mean fluorescence intensity (MFI) per islet in confocal images of human pancreas sections from non-diabetic (n = 21 islets, 6 donors), type 2 diabetic (n = 24 islets, 8 donors), and type 1 diabetic donors (n = 24 islets, 8 donors). Background (BKGD) indicates average taurine MFI in acinar tissue outside of the islet. One-way ANOVA: ND vs. T2D (*P= 0.0430), ND vs. T1D (ns, P= 0.6667). Center line indicates the mean.

d. Human pancreas sections immunostained for taurine, insulin, and glucagon from a nondiabetic, type 2 diabetic and, type 1 diabetic donor. Left panels show insulin and glucagon channels, while right panels show taurine channel from the same image. Images are representative of the dataset plotted in panel c. Scale bars 50 µm.

e. Representative confocal image of a monolayer of human islet endocrine cells showing immunostaining for GAD65, insulin, and glucagon (left panel) and GAD65 alone (right panel). Images are representative of 3 human islet preparations. Scale bar 20 μ m.



Extended Data Fig. 2. VGAT expression is concentrated in delta cells of human islets and alpha cells of rat islets; GABA colocalizes with GAD65 and VGAT in synaptic vesicles in neurons. a-b. Islets in a non-diabetic human pancreas (a) and rat pancreas (b) immunostained for VGAT, insulin, glucagon, somatostatin, and pancreatic polypeptide. VGAT is absent in most beta cells but present in somatostatin producing human delta cells and glucagon producing rat alpha cells. Results are representative of the dataset plotted in Figure 1b. Scale bar 50 μm.

c. Rat hippocampal neuron immunostained for GABA and the GABA biosynthesizing enzyme GAD65. Scale bar 10 μ m. Right panels show higher magnification views of the boxed region showing channels for: (1) GABA only; (2) GAD65 only; (3) GABA and GAD65. GAD65 and GABA colocalize in vesicles. Results are representative of n = 3 rat neuron preparations. Scale bar 5 μ m.

d. Rat hippocampal neuron immunostained for GAD65 and the vesicular GABA transporter VGAT, which is present in synaptic vesicle membranes. Scale bar 10 μ m. Right panels show higher magnification views of the boxed region showing channels for: (1) GAD65 only; (2) VGAT only; (3) GAD65 and VGAT. GAD65 and VGAT colocalize in synaptic vesicles. Results are representative of n = 3 rat neuron preparations. Scale bar 5 μ m.

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Extended Data Fig. 3. Characterization of GABA biosensor cells for detecting GABA released from human islets.

a. Schematic and image of the GABA biosensor cell assay setup (left panel). Biosensor cells consist of CHO cells stably expressing the heteromeric GABA_B receptor (GABA_B R1b and GABA_B R2) and the G-protein a subunit, Gaqo5 to allow for GABA detection by intracellular Ca²⁺ mobilization ($[Ca^{2+}]_i$) (right panel). GABA biosensor cells are preloaded with the $[Ca^{2+}]_i$ indicator Fura-2 and plated on poly-d-lysine coated cover slips in a perfusion chamber. Individual islets are placed on top of this layer of biosensor cells and connected to a closed bath small volume imaging chamber to ensure linear solution flow and fast exchange.

b. Titration of exogenous GABA showing concentration-dependence of Ca^{2+} flux in GABA biosensor cells. The plot shows the average 340/380 Fura-2 ratio of n = 5 GABA biosensor cells in the same field of view. Mean ± SEM.

c. Effect of the selective GABA_B receptor antagonist CGP5584 on biosensor cell responses to exogenously applied GABA. n = 5 biosensor cells in the field of view. Mean \pm SEM.

d. Biosensor cell intracellular Ca²⁺ responses remain elevated during sustained (30 min) exposure to GABA (100 μ M shown). n = 5 GABA biosensor cells in the field of view. Mean \pm SEM.

e. GABA release from a human islet maintained in 3 mM glucose. n = 5 GABA biosensor cells located under or immediately downstream of the islet. This is a representative trace of experiments performed on 40 human islet preparations. Mean \pm SEM.

f. Biosensor cells have tonic responses to continuously applied GABA and phasic responses to GABA pulses released from islets. Periods of pulsatile GABA release measured from n = 22 human islet preparations, 3 islets per preparation (black circles) as shown in panel e. Calculated periods for biosensor cell responses to continuously applied GABA (gray circles) at 0.1, 1, 10, and 100 μ M GABA as shown in panel d. Center line indicates the mean. **g.** GABA release from a human islet maintained in 3 mM glucose without addition of inhibitors. n = 5 GABA biosensor cells located under or immediately downstream of the islet. This is a representative trace of experiments performed on 40 human islet preparations. Mean \pm SEM.

h. Effect of the selective GABA_B receptor antagonist CGP5584 on biosensor cell detection of GABA released from a single human islet. n = 5 GABA biosensor cells located under or immediately downstream of the islet. Trace is representative of 3 independent experiments with different human islet preparations. Mean \pm SEM.



Extended Data Fig. 4. GABA release does not depend on glucose but is activated by VRAC opening.

a-b. Titration of glucose concentrations from 0–25 mM has no effect on islet GABA release. HPLC quantification of GABA released from rat (a) and human (b) islets during 30 mins static incubation in KRBH of the indicated glucose concentrations. n = 4 samples of 100 islets. One-way ANOVA, P = 0.5563 rat (a), P = 0.2053 human (b), ns = not significant. Mean \pm SEM.

c. HPLC quantification of GABA released from human islets in 5.5 mM glucose (n = 4 samples of 100 human islets), 30 mM KCl (n = 3 samples of 100 human islets), or diazoxide (100 μ M) (n = 4 samples of 100 human islets). One-way ANOVA, *P* = 0.1511, ns = not significant. Mean ± SEM.

d. Effect of the GAT inhibitors SNAP5114 (50 μ M), NNC05–2090 (50 μ M), and NNC711 (10 μ M) on biosensor detection of GABA secretion from a human islet. GAT inhibitors were present throughout the shaded portion of trace. Results are representative of the data plotted in panel e.

e. Quantification of GABA release following treatment with GAT inhibitors. Box extends from 25^{th} to 75^{th} percentiles, center line represents the median, whiskers represent smallest to largest values. n = 3 islets, One-way ANOVA, 0–20 min vs. 20–40 min (**P*= 0.002), 0–20 min vs. 40–60 min (**P*= 0.9194), 0–20 min vs. 60–80 min (**P*= 0.0058).

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Extended Data Fig. 5. Allylglycine inhibition of beta cell GABA content and secretion. a-b. Validation of GABA antibody via immunostaining of paraformaldehyde-fixed rat hippocampal neurons (a) or rat islet cell monolayers (b) for GAD65, GABA, and insulin (not shown) without or with addition of soluble GABA to the primary antibody incubation buffer; or without or with preincubation of cells with allylglycine (10 mM) to inhibit GABA biosynthesis. Images are representative of 3 experimental replicates. Scale bars 20 μm. **c.** Immunostaining of paraformaldehyde-fixed rat islets cell monolayers for GABA, insulin, and GAD65, following allylglycine (10 mM) addition and removal. Images are representative of the dataset plotted in panel d.

d. Quantification of GABA mean fluorescence intensity (MFI) in rat islet cell monolayers in allylglycine timecourse experiments shown in panel c. n = 4 coverslips. Mean \pm SEM. **e.** HPLC analysis of GABA release from human islets during a 30 min addition of allylglycine (no pre-incubation). n = 3 samples of ~100 islets each. Statistical analysis by two-tailed *t*-test, *P = 0.0104. Mean \pm SEM.



Extended Data Fig. 6. Human islet single-cell RNA-seq for expression of genes of interest.

a. Expression of neurotransmitter transporter family genes (SLC6A). Mean \pm SEM. **b.** Expression of genes of interest reported in the literature as related to GABA or putative GABA membrane transporters. Mean \pm SEM. Data shown are from two datasets^{54,55}, but results agree with and are representative of three different curated human single-cell RNA-seq datasets analyzed^{54–56} (see also Figure 4).





a-b. Still image, kymographs, and average trace from timelapse videos of Fura-2 $[Ca^{2+}]_i$ signals in GABA biosensor cells in a perfusion flow field in 3 mM glucose isotonic KRBH exposed to (a) 0.1, 1, and 10 μ M GABA, (b) downstream from a wild type mouse islet, and (c) downstream from a β c-LRRC8A^{-/-} mouse islet. GABA (1 μ M) is added to (c) at 23 min. GABA-responsive cells were selected for analysis, while unresponsive cells were not analyzed. Data are representative of three independent experiments. See also Supplementary Videos 1–3.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

ACKNOWLEDGMENTS

This work was funded by the Intramural Research Program of UF's Wertheim College of Engineering and J. Crayton Pruitt Family Department of Biomedical Engineering (E.A.P.), the Intramural Research Program of EPFL's School of Life Sciences (S.B.), the Diabetes Research Institute Foundation (DRIF), NIH grants R56DK084321 (A.C.), R01DK084321 (A.C.), NIDDK-supported Human Islet Research Network (HIRN, RRID:SCR_014393; https://hirnetwork.org; UC4DK104208 (E.A.P.)), a JDRF award (31-2008-416) to the European Consortium for Islet Transplantation (ECIT) Islets for Basic Research Program, the NIH NIDDK-funded Integrated Islet Distribution Program (IIDP) at City of Hope, NIH Grant # 2UC4DK098085 and a JDRF-funded IIDP Islet Award Initiative (E.A.P.), a JDRF Faculty Transition Award (1-FAC-2017-367-A-N) (E.A.P.), a JDRF Advanced Postdoctoral Fellowship (3-APF-2014-208-A-N) (E.A.P.), a Whitaker International Program Postdoctoral Scholarship (E.A.P.), the Swedish Research Council (P.-O.B.), the Novo Nordisk Foundation (P.-O.B.), the Family Erling-Persson Foundation (P.-O.B.), and the Stichting af Jochnick Foundation (P.-O.B.), the NIH NIDDK 1R01DK 106009 (R.S.), and the American Diabetes Association (1-18-IBS-229) (R.S.), and the Canadian Institutes for Health Research (PJT-159741, PJT-148652) (H.G.).

The work with human pancreatic sections and islets was made possible by the Human Islet Cell Processing Facility at the Diabetes Research Institute (University of Miami), ECIT, and the JDRF funded Network for Pancreatic Organ Donors with Diabetes (nPOD). Organ Procurement Organizations (OPO) partnering with nPOD to provide research resources are listed at http://www.jdrfnpod.org/for-partners/npod-partners/.

We wish to extend our thanks to the following individuals: Drs. Domenico Bosco and Thierry Berney, University of Geneva, and Dr. Lorenzo Piemonti, San Raffaele Scientific Institute, Milano for human islets through ECIT; Dr. Clayton Mathews, University of Florida, and Rita Bottino, Institute of Cellular Therapeutics, Allegheny Health Network, Pittsburgh, Pennsylvania, for human islets through the nPOD Islet Isolation Program; Dr. Klemens Kaupmann, Novartis Institutes for BioMedical Research, Switzerland, for providing the genetically-modified GABA biosensor CHO cells; and Dr. Stephen D. Roper for conceptual input on the biosensor cell approach and for critically reading the manuscript; Drs. Jeffrey Hubbell and Melody Swartz, EPFL and University of Chicago, for support; Chad Rancourt, University of Florida for mouse colony management, Miriella Pasquier, Kevin Johnson, Brandon Benjamin, Daniel Garcia, Paulo Parente, Abel Arzu, Lily Barash, Margaret Formoso, Rafael Arrojo e Drigo, and Alejandro Tamayo for technical assistance.

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Figure 1. Cytosolic pools of GABA are depleted in type 1 and type 2 diabetic islets **a.** A human pancreatic islet from a non-diabetic donor immunostained for GABA, insulin, and glucagon. Image is representative of the dataset plotted in panel b. Scale bar 50 μm. Right panels show higher magnification. Scale bar 10 μm.

b. Quantification of GABA colocalization with insulin (n = 15 islets, 9 donors), glucagon (n = 15 islets, 8 donors), and somatostatin (n = 15 islets, 3 donors) in human islets. One-way ANOVA: insulin vs. glucagon (*P< 0.0001), insulin vs. somatostatin (*P= 0.0182). Center line indicates the mean.

c-f. Human islets immunostained for GABA, insulin, and glucagon from a non-diabetic (c), type 2 diabetic (d), type 1 diabetic (e), and a type 1 diabetic donor with residual beta cell mass (f). Images are representative of the dataset plotted in panel g. Scale bars 50 µm. **g.** Quantification of GABA mean fluorescence intensity (MFI) per human islet from non-diabetic (n = 15 islets, 9 donors), type 1 diabetic (n = 15 islets, 8 donors), and type 2 diabetic donors (n = 12 islets, 8 donors). Background (BKGD) indicates average GABA MFI in acinar tissue outside of the islet. One-way ANOVA: ND vs. T2D (**P*<0.0001), ND vs. T1D (**P*<0.0001). Center line indicates the mean.

h. High performance liquid chromatography (HPLC) quantification of GABA content for human islet preparations from non-diabetic (n = 8 donors), type 2 diabetic (n = 3 donors), and type 1 diabetic donors (n = 2 donors). One-way ANOVA: ND vs. T2D (*P= 0.0196), ND vs. T1D (*P= 0.0106). Center line indicates the mean.

i. Human islet from a non-diabetic donor immunostained for GAD65, insulin, and glucagon. Image is representative of the dataset plotted in 1j. Scale bar 50 μ m.

j. Quantification of GAD65 colocalization with insulin (n = 24 islets, 9 donors), glucagon (n = 24 islets, 9 donors), and somatostatin (n = 11 islets, 3 donors) in human islets. One-way ANOVA: insulin vs. glucagon (*P< 0.0001), insulin vs. somatostatin (*P= 0.0257). Center line indicates the mean.

k-m. Human islets immunostained for GAD65 from non-diabetic, type 2 diabetic, and type 1 diabetic donors. Images are representative of the dataset plotted in panel n. Scale bars 50 μ m.

n. Quantification of GAD65 mean fluorescence intensity (MFI) per human islet from nondiabetic (n = 23 islets, 9 donors), type 1 diabetic (n = 23, 8 donors), and type 2 diabetic donors (n = 24 islets, 8 donors). Background (BKGD) indicates average GAD65 MFI in acinar tissue outside of the islet. One-way ANOVA: ND vs T2D (*P= 0.0380), ND vs. T1D (ns, P= 0.9511), T2D vs. T1D (ns, P= 0.0817). Center line indicates the mean.



Figure 2. Subcellular localization suggests a non-vesicular GABA release mechanism in beta cells a. A human islet from a non-diabetic donor immunostained for VGAT, insulin, and somatostatin. Image is representative of the dataset plotted in panel b. Scale bar 50 µm. Right panels show higher magnification views. Scale bar 10 µm.

b. Expression of VGAT is strongest in human delta cells and rat alpha cells but rare in beta cells of either species. Human islets stained for VGAT and insulin (n = 8 islets, 3 donors), glucagon (n = 7 islets, 3 donors), and somatostatin (n = 5 islets, 3 donors). Rat islets stained for VGAT and insulin (n = 7 islets, 3 donors), glucagon (n = 6 islets, 3 donors), and

somatostatin (n = 6 islets, 3 donors). One-way ANOVA, insulin vs. somatostatin, human (*P < 0.0001), insulin vs. glucagon, rat (*P < 0.0001). Center line indicates the mean.

c. GABA is non-vesicular and cytosolic in almost all human beta cells. A VGAT^{NEG}

primary human beta cell immunostained for GABA, VGAT, and insulin (not shown). Image is representative of n = 3 human islet preparations, 3 samples per preparation. Scale bar 50 μ m. Right panels show higher magnification views. Scale bar 2 μ m.

d. A rare VGAT^{POS} primary human beta cell showing colocalization of GABA and VGAT in vesicular structures. Image is representative of n = 3 human islet preparations, 3 samples per preparation. Scale bar 50 µm. Bottom panels show higher magnification views. Scale bar 2 µm.

e. A primary human delta cell immunostained for GABA, VGAT, and somatostatin (not shown) showing colocalization of GABA and VGAT in vesicular structures. Image is representative of n = 3 human islet preparations, 3 samples per preparation. Scale bar 50 μ m. Bottom panels show higher magnification views. Scale bar 2 μ m.

f-g. GABA is present in the cytosol and does not colocalize with insulin or GAD65 in vesicular structures in primary human (f) or rat (g) beta cells immunostained for GABA, insulin, and GAD65. Images are representative of n = 3 islet preparations, 3 samples per preparation. Scale bar 10 μ m. Inset images show single-channel higher magnification views. Scale bar 2 μ m.

h. Colocalization analyses between GAD65 (green) and the synaptic vesicle markers synaptophysin, SV2C, and VGAT (red) or GABA (red) in primary human beta cells, primary rat beta cells, and primary rat hippocampal neurons. Images are representative of data plotted in 2i. Scale bar 2 µm.

i. Quantification of colocalization in human beta cells for GAD65 with markers of synaptic vesicles synaptophysin (n = 14), SV2C (n = 3), and VGAT (n = 10); in primary rat beta cells (n = 11, 10, and 10); and primary rat hippocampal neurons (n = 6, 11, and 11). While GAD65 positive vesicles in rat neurons colocalize with synaptic vesicle markers, colocalization is rare or absent in human and rat beta cells. human beta cells:. Two-way ANOVA: human beta cells vs rat neurons for all markers (*P< 0.0001), rat beta cells vs. rat neurons for all markers (*P< 0.0001).

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Figure 3. Islet GABA secretion is pulsatile and depends on GABA content

a. GABA release from a human islet maintained in 3 mM glucose, detected by cytosolic Ca^{2+} flux in GABA_B receptor-expressing biosensor cells (black trace). Biosensor cell responses to no islets or to GABA only are shown for comparison (blue traces). Unless otherwise specified, in this and all subsequent Figures, the plot shows the average 340/380 Fura-2 ratio from 5 GABA biosensor cells located under the islet, hereafter labeled as GABA release. This is a representative trace of experiments performed on n = 40 human islet preparations.

b. The GABA_B receptor antagonist CGP55845 (10 μ M) blocks biosensor cell detection of GABA pulses from a single human islet. Results are representative of n = 3 human islet preparations, 3 islets per preparation.

c. Periods of pulsatile GABA release measured from n = 22 human islet preparations, 3 islets per preparation. Mean \pm SEM.

d. Representative trace of GABA release pulses from a human islet at different glucose concentrations (1G = 1 mM, 3G = 3 mM, 11G = 11 mM). Results are representative of data plotted in panel e.

e. Quantification of traces shows no difference in the amount of released GABA per pulse at different glucose concentration (n = 3 islet preparations, 18–20 islets per each preparation). One-way ANOVA: 1G vs. 3G (ns, P= 0.6181), 1G vs. 11G (ns, P= 0.1684), 3G vs. 11G (ns, P= 0.5302). Mean ± SEM.

f. Insulin and GABA released during human islet perifusion. GABA and insulin were quantified in the same sample by HPLC and ELISA (n = 3 samples of 100 islets each). One-way ANOVA: for insulin release, 3G vs. 16.7G (*P= 0.0046), 3G vs. KCl (*P= 0.0007); for GABA release 3G vs. 16.7G (ns. P= 0.5034). 3G vs. KCl (ns. P= 0.2341)

GABA release, 3G vs. 16.7G (ns, P = 0.5034), 3G vs. KCl (ns, P = 0.2341). **g.** Islet GABA secretory pulses before and during (shaded area) KCl depolarization (30 mM). Trace is representative of n = 3 human islet preparations, 3 islets per preparation. **h.** Islet GABA secretory pulses in the presence and absence (shaded area) of nominal Ca²⁺. Trace is representative of n = 3 human islet preparations, 3 islets per preparation. **i.** Quantification of GABA release during KCl depolarization and absence of nominal Ca²⁺ compared to control (n = 15 islets, 3 donors). Two tailed *t*-test: KCl vs. hypothetical mean of 1.0 (*P= 0.0069), 0 Ca²⁺ vs. hypothetical mean of 1.0 (ns, P= 0.1202). Mean ± SEM. **j.** HPLC quantification of GABA release from human islets in the presence and absence of nominal Ca²⁺ (n = 4 samples of 100 islets each). One-way ANOVA: control vs. 0 Ca²⁺ (ns, P= 0.5231), control vs. 0 Ca²⁺ + thapsi (ns, P= 0.2946), 0 Ca²⁺ vs. 0 Ca²⁺ + thapsi (ns, P= 0.8851). Mean ± SEM.

k. Trace of GABA release from a human islet before (black trace) and after (red trace) applying the GAD65 inhibitor allylglycine (10 mM). Trace is representative of n = 3 islet preparations, 3 islets per preparation.

I. Trace of GABA release from a human islet before (black trace) and after (green trace) applying the GABA transaminase inhibitor γ -vinyl GABA (10 μ M). Trace is representative of n = 3 islet preparations, 3 islets per preparation.

m. Quantification of GABA release from human islets treated with allylglycine or γ -vinyl GABA. (n = 3 islet preparations, 4 islets per preparation). Two-tailed student's *t*-test: control vs. allylglycine (**P*=0.0353), control vs. γ -vinyl GABA (**P*=0.0411). Mean \pm SEM.

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b. Gene expression for SLC6A and other proteins of interest for GABA transport and biosynthesis in human beta cells from an analysis of a curated human islet single-cell RNA-seq dataset (n = 158 cells)⁵⁶. Mean \pm SEM. Results are representative of three different human single-cell RNA-seq datasets analyzed (Extended Data 6 and Supplementary Table 1).

c. Representative confocal image of a human pancreas section immunostained for TauT and insulin. Scale bar 50 μ m. Right panels show higher magnification views of the boxed region with individual channels for (1) TauT and (2) insulin. Image is representative of n = 3 donors, 3 islets per donor. Scale bar 20 μ m.

d. Representative confocal microscopy images of an islet in a human pancreas section immunostained for LAT2 (upper panel) and 4F2HC (lower panel). Scale bar 50 μ m. Right panels show higher magnification views of the boxed region showing individual channels for (1) LAT2 and (2) 4F2HC. Image is representative of n = 3 donors, 3 islets per donor. Scale bar 20 μ m.

e. Uptake of radiolabeled 3H-GABA by human islets with competitive inhibition from unlabeled GABA, taurine, beta-alanine, leucine, or glycine; or blocking the GABA transporters with a mixture of SNAP5114 (50 μM), NNC05–2090 (50 μM), and NNC711 (10 μM) (GAT blockers) (n = 3 samples of 500 islets each). One-way ANOVA: control vs. GABA (*P< 0.0001), control vs. taurine (*P< 0.0001), control vs. β-alanine (*P< 0.0001), control vs. GAT blockers (*P< 0.0001), control vs. leucine (*P< 0.0001), control vs. GAT blockers (*P< 0.0001), control vs. leucine (*P= 0.0139). Data are representative of 3 islet preparations (2 human, 1 rat). Center line indicates the mean.

f. HPLC measurement of cumulative GABA and taurine release from human islets in KRBH buffer of varying osmolarity (n = 3 samples, 100 islets each). One-way ANOVA: for GABA, 300 vs. 160 mOsm/L (*P< 0.0001), 300 vs. 250 mOsm/L (*P= 0.0288). Data are representative of 4 human islet preparations. Center line indicates the mean.

g. Western blot for LRRC8A and β -actin in WT and LRRC8A^{-/-} MIN6 insulinoma cells. Results are representative of 3 independent immunoblots with similar results.

h. HPLC measurements of rate of GABA release from wild type (WT) or LRRC8A^{-/-} MIN6 insulinoma cells during static incubation in 3 mM glucose isotonic (300 mOsm/L) or hypotonic (250 mOsm/L) KRBH buffer (n = 3 samples of 40,000 cells each). WT and LRRC8A^{-/-} MIN6 cells were transfected with a plasmid encoding human GAD65 24 hours prior to the experiment as they do not endogenously express either GAD65 or GAD67. Twoway ANOVA: WT iso vs. WT hypo (**P*< 0.0001), WT iso vs. LRRC8A^{-/-} iso (**P*= 0.0053), LRRC8A^{-/-} iso vs. LRRC8A^{-/-} hypo (ns, *P*= 0.2974). Center line indicates the mean.

i. Western blot for LRRC8A in mouse islets isolated from WT (LRRC8A^{+/-} littermates heterozygous for floxed allele) and beta cell-specific knockout (β c-LRRC8A^{-/-}) mice. Western blot was performed once using islets pooled from 2 donors of each genotype. **j.** HPLC measurements of cumulative GABA release from WT and β c-LRRC8A^{-/-} mouse islets in 3 mM glucose isotonic (300 mOsm/L) or hypotonic (250 mOsm/L) KRBH buffer. n = 3 samples of 20 islets each. Two-way ANOVA: WT iso vs. WT hypo (*P< 0.0001), β c-LRRC8A^{-/-} iso vs. β c-LRRC8A^{-/-} hypo (ns, P= 0.8057). Mean ± SEM.

k. Biosensor cell traces of GABA release from WT and β c-LRRC8A^{-/-} mouse islets during isotonic and prolonged (10 min+) hypotonic stimulation. Islets used for these experiments were confirmed for loss of LRRC8A by Western blot (panel g). Results are representative of data plotted in panel 1.

l. GABA release detected by biosensor cells from WT (n = 5) and β c-LRRC8A^{-/-} (n = 3) mouse islets. Data are representative of two independent islet isolation. Two-way ANOVA of

log transformed data: WT iso vs. WT hypo (*P= 0.0003), WT iso vs. β c-LRRC8A^{-/-} iso (*P= 0.0003), β c-LRRC8A^{-/-} iso vs. β c-LRRC8A^{-/-} hypo (ns, P= 0.9999). Mean ± SEM. **m.** Western blot for LRRC8A in human islets infected with Ad-mCherry-Scramble-shRNA or Ad-mCherry-hLRRC8A-shRNA adenovirus. Western blot was performed once. **n.** Knockdown of LRRC8A in human islets reduces basal GABA pulses. Results are

representative of the data plotted in panel o.

o. Quantification of biosensor cell detection of GABA release from human islets infected with Ad-mCherry-Scramble-shRNA (n = 3 islets) or Ad-mCherry-hLRRC8A-shRNA (n = 6 islets). Data are representative of two independent human islet preparations. Two-tailed *t*-test (*P= 0.0065). Mean ± SEM.

p. HPLC measurement of dynamic GABA release from nondiabetic human islets during isotonic (300 mOsm/L) and prolonged hypotonic (250 mOsm/L) stimulation. n = 3 samples of 100 islets each. Mean \pm SEM.





a. Human islet perifusion in 5 mM glucose showing reversible inhibition of insulin secretion in response to exogenous GABA (10 μ M). n = 3 islet preparations. Mean \pm SEM.

b. Human islet perifusion showing differential glucose-stimulated insulin response in islets treated with GAD65 inhibitor allylglycine (10 mM, red trace) vs. control (black trace). n = 3 islet preparations. Mean \pm SEM.

c. Quantification from islet perifusion during 11 mM glucose stimulation phase, control (black trace), with allylglycine (red trace). n = 3 islet preparations (average of > 3 islets per preparation), Two-tailed *t*-test, *P = 0.0175. Mean \pm SEM.

d. Four independent traces of cytosolic Ca²⁺ levels measured by Fluo-4 dye in human islets during successive addition of 3 mM glucose, 16.7 mM glucose, and GABA (10 μ M). GABA induces a rapid drop of cytosolic Ca²⁺ that desensitizes after ~90s. Data are representative of three human islet preparations.

e. Quantification of Ca²⁺ levels in human islets under 16.7 mM glucose stimulation during the first 2 minutes after addition of GABA. n = 4 islets. Two-tailed *t*-test, **P*= 0.0132. Mean \pm SEM.

f. Effect of GAD65 inhibitor allylglycine (10 mM) on serotonin/insulin secretion from human islets at 3 mM glucose concentration using serotonin biosensor cells. Trace is representative of experiments performed in n = 3 human islet preparations, 3 islets/ preparation.

g. Effect of the GABA transaminase blocker γ -vinyl GABA (10 μ M) on serotonin/insulin secretion from human islets at 3 mM glucose concentration. Ca²⁺ trace is representative of experiments performed in n = 3 human islet preparations, 3 islets/preparation.

h. Quantification of serotonin/insulin secretion from human islets treated with allylglycine or γ -vinyl GABA from experiments performed. n = 3 human islet preparations, 3 islets/ preparation. One-way ANOVA, **P*<0.0001. Mean ± SEM.

i. Periodicity of serotonin/insulin secretion from human islets (n = 28 biosensor cells) and the increase in periodicity variance in the presence of the GABA_A receptor antagonist SR 95531 (10 μ M) (n = 35 biosensor cells). Data are pooled from 3 independent human islet preparations, > 3 islets/preparation. Two-sided F test to compare variances, **P* = 0.0002. Mean ± SEM.

j. Traces of the sum of fluorescent exocytotic signals in three individual beta cells from different islet regions showing periodic, synchronous exocytosis at 16 mM glucose (left) and asynchronous, diminished exocytosis after adding the GABA_A receptor agonist muscimol (100 μ M, right). Traces are representative of n = 3 islet preparations, 3 islets per preparation.

k. Quantification of the amplitude of beta cell fluorescent exocytotic signals in islets treated with muscimol. n = 3 human islet preparations, 3 islets/preparation, all measurements depicted. Two-tailed *t*-test, **P* < 0.0001. Mean ± SEM.



Figure 6. Cytosolic GABA secretion is interrupted in human islets from type 2 diabetic donors **a.** Absence of GABA secretion detected by biosensor cells from an islet taken from a type 2 diabetic donor (black trace), but distinct GABA secretion is detected from an islet from a non-diabetic donor (gray trace). Addition of exogenous GABA to a type 2 diabetic islet induces a strong response from biosensor cells. Representative of experiments performed in 5 islet preparations from type 2 diabetic donors and 40 non-diabetic donor preparations, 3 islets/preparation.

b. Quantification of GABA release from non-diabetic (n = 4) and type 2 diabetic (n = 5) human islets. Two-tailed *t*-test, *P = 0.0418. Mean \pm SEM.

c. Ca²⁺ flux in diabetic human islets during stimulation with KCl. Representative of experiments performed in 5 islet preparations from diabetic donors.

d. Pulsatile GABA release from an islet from a type 2 diabetic donor before (black trace) and after (green trace) exposure to the GABA transaminase inhibitor γ -vinyl GABA (10 μ M, 1 hour). Representative of n = 5 islets.

e. Quantification of GABA release pulse amplitude from type 2 diabetic human islets before and after exposure to γ -vinyl GABA. n = 3 human islet preparations (average of 3 islets/ preparation). Two-tailed *t*-test, **P* = 0.0257. Mean ± SEM.

f. Insulin/serotonin release detected by serotonin biosensor cells from an islet from a type 2 diabetic donor before (black trace) and after (green trace) exposure to γ -vinyl GABA (10 μ M, 1 hour). Representative of n = 5 islets.

g. Periods of insulin/serotonin release from islets from a type 2 diabetic donor before (black dots) and after (green dots) exposure to γ -vinyl GABA (10 μ M, 1 hour) from experiments performed as in (f). n = 5 islets. Two-tailed *t*-test to compare means, **P* = 0.01, two-sided F test to compare variances, **P* = 0.0118. Mean ± SEM.

h-i. Confocal images of islets from non-diabetic (h) and type 2 diabetic donors (i) immunostained for GAD65 and Golgi resident protein GCP60. Arrows point at GAD65 accumulation in Golgi membranes. Images are representative of data plotted in panel j. Scale bar 10 μ m.

j. Quantification of GAD65 immunostaining intensities in Golgi membranes in beta cells of pancreata from non-diabetic and type 2 diabetic donors. GAD65 staining in Golgi membranes is expressed relative to the total cellular GAD65 staining (GAD65). n = 18 islets from 3 donors per group. Two-tailed *t*-test, **P* < 0.0001. Mean ± SEM.