GIPR Is Predominantly Localized to Nonadipocyte Cell Types Within White Adipose Tissue

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The incretin hormone glucose-dependent insulinotropic polypeptide (GIP) augments glucose-dependent insulin secretion through its receptor expressed on islet β cells. GIP also acts on adipose tissue; yet paradoxically, both enhanced and reduced GIP receptor (GIPR) signaling reduce adipose tissue mass and attenuate weight gain in response to nutrient excess. Moreover, the precise cellular localization of GIPR expression within white adipose tissue (WAT) remains uncertain. We used mouse genetics to target Gipr expression within adipocytes. Surprisingly, targeting Cre expression to adipocytes using the adiponectin (Adipoq) promoter did not produce meaningful reduction of WAT Gipr expression in Adipoq-Cre:Gipr^{flx/flx} mice. In contrast, adenoviral expression of Cre under the control of the cytomegalovirus promoter, or transgenic expression of Cre using nonadipocyte-selective promoters (Ap2/Fabp4 and Ubc) markedly attenuated WAT Gipr expression. Analvsis of single-nucleus RNA-sequencing, adipose tissue data sets localized Gipr/GIPR expression predominantly to pericytes and mesothelial cells rather than to adipocytes. Together, these observations reveal that adipocytes are not the major GIPR⁺ cell type within WAT-findings with mechanistic implications for understanding how GIP and GIP-based co-agonists control adipose tissue biology.

Incretin hormones are produced in specialized enteroendocrine cells and amplify meal-stimulated insulin release

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following food ingestion. The two incretins, glucosedependent insulinotropic polypeptide (GIP) and glucagonlike peptide-1 (GLP-1), exert their actions through structurally related yet distinct receptors (1). The major target for incretin action is the islet β cell, wherein GIP and GLP-1 potentiate insulin secretion through cyclic AMPdependent and -independent pathways (1).

The molecular cloning of incretin receptor cDNAs enabled identification of extrapancreatic expression of both incretin receptors, consistent with characterization of multiple actions for GIP and GLP-1 beyond the β cell (1,2). GLP-1 receptor mRNA transcripts have been identified within the central and enteric nervous systems, the heart, gastrointestinal tract, kidney, blood vessels, and immune cells (3–5), lending support for multiple direct actions of GLP-1 in peripheral tissues. Similarly, the tissue distribution of GIP receptor (GIPR) mRNA transcripts includes the brain, heart, gastrointestinal tract, blood vessels, and adipose tissue (6).

Among key differences in the extrapancreatic biology of GIP but not GLP-1 is that the former acts directly on adipose tissue (1,7). These actions of GIP have been studied using differentiated adipose tissue cell lines, primary adipocyte cultures, and experiments with animals and humans (8–14). Within white adipose tissue (WAT), GIP promotes both lipolysis and lipid accretion, glucose uptake, insulin sensitization, and adipokine expression (12,13,15–20). GIP

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also modifies adipose tissue biology through actions on blood vessels, and immune cells. For instance, GIP rapidly augments adipose tissue blood flow (21) and either augments or suppresses WAT inflammation through direct actions on immune cells (13,18,22,23).

Interpreting the actions of GIP on adipose tissue has been complicated by paradoxical observations that both sustained GIPR agonism and attenuation of GIPR signaling produce overlapping phenotypes in animals, including reduction of WAT inflammation, reduced WAT mass, resistance to weight gain, and improvement of insulin sensitivity (7,18,24–31). Understanding how gain or loss of GIP actions within unique adipose tissue GIPR⁺ cell types has been hampered in part by 1) the lack of highly specific validated antisera or labeled analogs for detection of the GIPR (32,33), and 2) a paucity of in situ hybridization or single-cell RNA sequencing (scRNA-seq) data for detection of the WAT GIPR. Hence, the cellular localization of GIPR in different adipose tissue depots remains incompletely understood.

In view of multiple studies linking expression of GIPR to adipocytes, we attempted to understand the metabolic consequences of targeting the adipocyte *Gipr* using mouse genetics. Surprisingly, although expression of Cre under control of the well-characterized adiponectin promoter recombined Gipr genomic DNA in WAT, we did not observe meaningful reduction of Gipr expression in WAT depots of Adipoq-Cre:Giprfix/fix mice. In contrast, expression of Cre recombinase under the control of the human adipocyte fatty acid binding protein (Ap2) Fabp promoter, or using *Ubc*-CRE^{ERT2} to direct widespread Cre expression, resulted in marked reduction of WAT Gipr mRNA transcripts in multiple WAT depots. Consistent with these findings, Gipr-directed reporter expression was not detected within the majority of adipocytes analyzed using a Gipr-Cre mouse to identify transcriptional domains of endogenous Gipr promoter activity. Finally, publicly available single-cell RNA-seq (scRNA-seq) data identified WAT Gipr/GIPR expression predominantly within pericytes and mesothelial cells (34). Taken together, these findings refine our understanding of WAT Gipr expression, providing insights for guiding interpretation of data linking GIP action to changes in adipose tissue biology.

RESEARCH DESIGN AND METHODS

Animal Models

Animals were obtained from The Jackson Laboratory (Bar Harbor, ME). *Gipr*^{flx/flx} (35) *Adipoq*-Cre (36), *Adipoq*BAC-Cre (catalog 028020 (37)), *Mip*-Cre^{ERT} (catalog 024709 (35,38), α *MHC*-Cre^{ERT} (32), *Ubc*-Cre^{ERT2} (catalog 008085 (39)), *Adipoq*BAC-Cre^{ERT} (catalog 024671 (40)), and *Fabp4*(*Ap2*)-Cre (catalog 005069 (41)) mice have been previously described. *Gipr*-Cre (knock-in) mice were generated using CRISPR/Cas9, as previously described (42) and bred with Rosa26-LacZ (B6;129S4-*Gt*(*ROSA*)*26Sort*^{m1Sor}/*J*/*J* (catalog 003309) or Rosa29-mTmG (*Gt*(*ROSA*)*26Sort*^{m4(ACTB-tdTomato,-EGFP)Luo/J} (catalog 007576) reporter mice. Male mice were used for all studies.

PCR Analysis

RNA isolation and quantitative PCR were carried out as previously described (35). PCR of genomic DNA was done using PrimeSTAR GXL DNA Polymerase (Takara Bio; catalog R050A). Primer sequences are described in Supplementary Table 1.

In Vivo Studies

All animal studies were conducted under protocols approved by the Animal Care Committees of the Toronto Centre for Phenogenomics, the Duke Molecular Physiology Department, and the University of Cambridge Animal Welfare and Ethical Review Body and conformed to the Animals (Scientific Procedures) Act 1986 Amendment Regulations (SI 2012/3039). Briefly, intraperitoneal and oral glucose tolerance tests (1.5 g/kg glucose) were performed in animals after a 5-h fast. For intraperitoneal glucose tolerance tests, animals were given PBS or D-Ala2 GIP (4 nmol/kg; Chi Scientific) intraperitoneally 10 min prior to glucose administration. Insulin tolerance was measured following administration of 0.7 units/kg fast-acting insulin (Humalog) in mice fasted for 5 h. The high-fat diet (HFD) feeding was a 45% fat diet from Research Diets (D12451). Body composition was measured with an EchoMRI device.

In Vivo Adenovirus Treatment

An adenoviral vector containing the human adenovirus type5 (dE1/E3) viral backbone encoding both Cre recombinase and GFP through separate cytomegalovirus (CMV) promoters was gifted by Dr. Andras Nagy (Lunenfeld-Tanenbaum Research Institute, Mt. Sinai Hospital, Toronto, ON, Canada). The virus was used at a titer of 1×10^{10} PFU/mL and a total of 50 µL was injected directly into a single inguinal fat pad in mice lightly anesthetized with isoflurane. Virus containing an empty vector was injected at the same amounts into the contralateral inguinal fat pads were harvested, flash frozen, and stored at -80C until used for RNA analysis.

HFD Feeding of Ap2-Cre Mice

Littermate, age-matched controls were weened at 3–4 weeks of age and maintained on standard rodent chow until 8 weeks of age. The diet was then switched to a 45% fat diet and the mice were maintained on this diet until study termination.

Whole-Mount β-Galactosidase Assay

Tissues were harvested from 11-month-old male mice that were hemizygous for *Gipr*-Cre and heterozygous for *ROSA*26-LacZ or heterozygous for *ROSA*26-LacZ (negative control). Tissues were rinsed in PBS and transferred to 6-well plates, where they were fixed for 2 h (in calcium- and

magnesium-free PBS containing 1% paraformaldehyde, 0.2% glutaraldehyde, and 0.02% Nonidet P-40) at 4°C using an orbital shaker. Samples were then washed twice (20 min each) in PBS and incubated in the dark overnight (16 h) at 37°C in β -galactosidase substrate (calcium- and magnesium-free PBS containing 5 mM potassium ferricyanide, 5 mmol/L potassium ferrocyanide, 2 mmol/L magnesium chloride, 0.02% NP-40, 0.01% sodium deoxycholate, and 1 mg/mL X-gal substrate). The following day, samples were rinsed twice in PBS as described, fixed in 10% neutral buffered formalin overnight at 4°C, and transferred to 70% ethanol until imaging. Whole-mount tissues were imaged using a Leica MZ6 stereomicroscope with an attached MC170 HD digital camera (Leica Microsystems Inc., Concord, ON, Canada).

Whole-Mount Confocal Microscopy

Adipose tissues were harvested from 14-week-old female mice that were hemizygous for Gipr-Cre and heterozygous for mTmG or heterozygous for mTmG (negative control), rinsed with PBS, cut into 0.5- to 1-cm pieces, transferred to 12-well plates, and fixed in 1% paraformaldehyde for 1 h at room temperature. Tissues were then washed three times (10 min each) in PBS containing 0.3% Triton X-100, followed by an additional three washes (10 min each) in PBS and then incubated in Lipidtox Deep Red (1:1,000 in PBS, catalog H34477; Thermo Fisher Scientific, Mississauga, ON, Canada) for 30 min at room temperature. Tissues were placed on a glass coverslip and saturated with DAPI-containing mounting media (Vectashield, catalog H-1200; Vector Laboratories Inc., Burlington, ON, Canada) and imaged using an inverted confocal laser microscope.

Single-Nucleus RNA-Seq Analysis

Detection of *Gipr/GIPR* and other class B, G protein-coupled receptor (GPCR) mRNA transcripts within mouse and human adipose tissue depots was accomplished using publicly available data derived from single-nucleus RNAseq analyses, as previously described (34), accessed from the Broad Institute single-cell portal (https://singlecell. broadinstitute.org/single_cell).

Statistics

Data are presented as mean \pm SEM. Differences were determined by Students *t* test, one- or two-way ANOVA, as appropriate. A Tukey test was used for post hoc analysis of ANOVAs, where appropriate. *P* < 0.05 was set as the criterion for statistically significant difference.

Data and Resource Availability

The original data in this article are available upon reasonable request from the authors. The *Gipr* floxed mice are available for collaborative sharing from the authors (contact D.J.D.). All of the other mouse lines and reagents described herein are available from commercial sources.

RESULTS

Gipr mRNA Transcripts Are Not Reduced in Adipose Tissues From *Adipoq-Cre:Gipr^{fix/fix}* Mice

Several studies have reported targeting of murine *Gipr* expression in WAT; however, the extent of *Gipr* knock-down within adipocytes in vivo was not described (41,43). Nevertheless, based on reports localizing *Gipr* expression to adipocytes (43–45), we sought to inactivate adipocyte *Gipr* expression using the widely used adiponectin-Cre system (46). Accordingly, we generated *Adipoq-Cre:Gipr*^{flx/flx} mice by crossing *Gipr*^{flx/flx} mice (35,47) with mice expressing *Cre* driven by the *Adipoq* promoter (36). Surprisingly, levels of *Gipr* mRNA transcripts in WAT or brown adipose tissue (BAT) from *Adipoq-Cre:Gipr*^{flx/flx} mice were not reduced (Fig. 1*A*).

We next generated a second mouse model using an independently generated Adipoq-Cre mouse that uses a BAC transgene containing the majority of the Adipoq regulatory elements (AdipoqBAC-Cre) (37), a mouse line successfully used by multiple groups to achieve adipocyte-selective gene recombination (48), including our own laboratory (49). Unexpectedly, AdipoqBAC- $Cre:Gipr^{flx/flx}$ mice also failed to exhibit reduced Gipr expression in WAT and BAT (Fig. 1B and C) despite expressing Cre and Adipoq at levels similar to wild-type (WT) and AdipoqBAC-Cre control mice (Fig. 1B). Collectively these findings indirectly imply that the majority of adiponectin⁺ adipocytes do not express the Gipr within adipose tissue in vivo.

We previously achieved reduction of *Gipr* expression, using the same *Gipr*^{flx/flx} mice, in β cells (35), cardiomyocytes (32), and BAT (47) using *Mip-Cre*^{ERT}, $\alpha MHC-Cre^{ERT}$, and *Myf5-Cre* mice, respectively. To explain the lack of *Gipr* knockdown in WAT, we examined relative *Cre* expression across different mouse models. WAT *Cre* expression driven by the *AdipoqBAC* promoter was comparable to levels seen in primary islets, driven by *Mip*, but lower than levels detected in the heart, driven by αMHC . (Fig. 1*D*). *Gipr* mRNA transcript levels were reduced in heart and islet tissues of αMHC -*Cre:Gipr*^{flx/flx} and *Mip-Cre:Gipr*^{flx/flx} mice, respectively (Fig. 1*D*), consistent with previous observations (32,35). In contrast, *Gipr* mRNA transcripts were not reduced in adipose tissue depots from *AdipoqBAC-Cre:-Gipr*^{flx/flx} mice (Fig. 1*D*).

Gipr Expression Can Be Reduced in WAT of *Gipr^{fix/fix}* Mice Through Nonadipocyte Selective *Cre* Expression

The failure to knock down WAT *Gipr* mRNA transcript levels using *Adipoq*-Cre raised several possibilities. First, we hypothesized that the adipocyte *Gipr* gene may be uniquely inaccessible to Cre within adipocytes, thereby preventing Cre-mediated recombination of the floxed alleles. Alternatively, we surmised that cellular *Gipr* expression may be inversely correlated to *Adipoq* expression, implying that the *Gipr* may not be expressed in the majority of white adipocytes. To examine these possibilities, we injected a *Cre*-expressing adenovirus directly into the inguinal adipose



Figure 1 – Cre recombinase expression driven by the adiponectin promoter does not reduce *Gipr* mRNA transcripts in adipose tissue. *A*) *Gipr* expression in epididymal WAT and interscapular BAT. Cre expression is driven by transgenic expression of *Adipoq-Cre* (36). (n = 3-8). *B*) *Gipr, Cre,* and *Adipoq* expression in epididymal WAT and interscapular BAT. For reference, the average cycle threshold values for *Gipr* and *Adipoq* were 26.28 and 18.77, respectively, in WAT and 28.13 and 20.73, respectively, in BAT. *Cre* expression is driven by the *AdipoqBAC* promoter (*AdipoqBAC-Cre* (37)). (n = 3). *C*) *Gipr* expression in various adipose tissue depots in mice with the *AdipoqBAC-Cre* transgene. (n = 3). *D*) *Cre* and *Gipr* expression in tissues, including epididymal WAT from various Cre recombinase models crossed with *Gipr*^{fix/fix} mice. (n = 2-3). *A*–*C*) For relative RNA expression values, the values are normalized to expression in the Cre controls (*Adipoq-Cre* in *A*; *AdipoqBAC-Cre* in *B* and *C*). *D*) Values are normalized to levels for *MIP-Cre*^{ERT}. *P < 0.05 vs control. ND = Not detected.



Figure 2—Complementary genetic strategies enable reduction of *Gipr* expression in adipose tissue. *A*) An adenoviral vector expressing *Cre* and *Gfp* under control of the CMV promoter was injected into an inguinal adipose depot of *Gipr*^{fix/fix} mice; empty vector was administered into the contralateral depot as a control. Quantitative PCR was used to determine the expression of *Gfp* and *Gipr* in both depots. (n = 3 mice.) *B*) Blood glucose levels during an intraperitoneal glucose tolerance test in which PBS or GIP (4 nmol/kg) was administered 10 min before glucose in control (*Ubc-Cre*^{ERT2}; n = 8-13) or *Ubc-Cre*^{ERT2}:*Gipr*^{fix/fix} (n = 7-9) mice. *C*) *Gipr* expression in primary islets, whole-brain, and epididymal adipose tissue. (n = 4). *D*) *Gipr* expression in epididymal adipose tissue from *Adipoq-Cre*^{ERT} (control) and *Adipoq-Cre*^{ERT}:*Gipr*^{fix/fix} mice harvested 2 weeks following tamoxifen treatment, when mice were 10 weeks of age. (n = 4). *E*) PCR analysis of genomic DNA (gDNA) from epididymal WAT from different genetic mouse models. Nonrecombined DNA produces an 850-bp product. For relative mRNA values, expression in panel *A* was normalized to control; for panel *C*, values were normalized relative to *Ubc-Cre*^{ERT}, **P* < 0.05 vs control. AUC, area under the curve; KO, knockout; NTC, no template control.

depot of $Gipr^{fix/flx}$ mice as a means of producing widespread expression of Cre within WAT (50). Mice injected with *Ad-CMV(GFP)-Cre* exhibited a 70% knockdown in inguinal adipose tissue *Gipr* expression (Fig. 2A), illustrating the susceptibility of the *Gipr* allele within WAT of *Gipr*^{fix/fix} mice to recombination and subsequent reduction of WAT *Gipr* mRNA transcripts.

To obtain complementary evidence supporting these observations, we used $Ubc-Cre^{ERT2}$ mice, which express Cre under the control of the human ubiquitin C promoter in most cell types (39). Consistent with loss of the insulinstimulating actions of GIP in $Gipr^{\beta}$ cell-/- mice (35), tamoxifen-treated $Ubc-Cre^{ERT2}$: $Gipr^{fbx/fbx}$ mice failed to exhibit reduction of glucose levels in response to exogenous GIP (Fig. 2B). Importantly, levels of Gipr mRNA transcripts were markedly reduced in the islets, brain, and WAT of $Ubc-Cre^{ERT2}$: $Gipr^{fbx/fbx}$ mice (Fig. 2C). Thus, both $UbcCre^{ERT2}$ and

Ad-CMV-*Cre* can drive *Cre* expression enabling recombination of the *Gipr* gene and reduction of *Gipr* mRNA transcripts in WAT from $Gipr^{flx/flx}$ mice.

AdipoqBAC-Cre Mice Exhibit Recombination of the Genomic Gipr Locus in WAT, Without Reduced Adipocyte Gipr Expression

 $Ubc-Cre^{ERT2}$ mice require tamoxifen to induce Cre recombinase activity, whereas the AdipoqBAC-Cre mice (Fig. 1B–D) exhibit constitutive expression of Cre recombinase, including during development. We wondered whether postnatal induction of Cre expression may confer preferential recombination of floxed adipocyte Gipr alleles. To assess this possibility, we crossed Gipr^{flx/flx} mice with AdipoqBAC-Cre^{ERT} mice (40), which require tamoxifen for conditional induction of Cre activity. Following the same tamoxifen protocol used for the Ubc-Cre^{ERT2} model (Fig.

2B and C), treatment of AdipogBAC-Cre^{ERT}:Gipr^{fix/fix} mice with tamoxifen at 8 weeks of age failed to reduce epididymal Gipr expression (Fig. 2D). However, PCR analysis of epididymal adipose tissue DNA using primers that span both loxP sites demonstrated that the Cre recombinase effectively induced recombination of the Gipr genomic DNA (Fig. 2E). In control adipose tissue samples (WT or *Gipr*^{flx/flx}), only the full-length genomic DNA PCR product was amplified (no recombination). In contrast, both the full-length and Cre-generated products were amplified in genomic DNA from AdipoqBAC-Cre^{ERT}:Gipr^{fix/fix} adipose tissue, and only the truncated Cre-generated PCR product was amplified in genomic DNA from Ubc-Cre^{ERT}:Gipr^{fix/fix} adipose tissue (Fig. 2E). Thus, the Adipoq promoter is capable of generating sufficient Cre expression to permit recombination of the Gipr allele yet does not alter Gipr mRNA levels within WAT.

Ap2-Cre:Gipr^{flx/flx} Mice Exhibit Reduced Gipr Expression in Adipose Tissue and Brain

Several metabolic phenotypes were described for mice with Gipr knockdown in adipose tissue, generated using Ap2-Cre mice (41), including a modest reduction in body weight after HFD feeding, together with improved glucose tolerance, and reduced hepatic steatosis. Intriguingly, the reductions in body weight were driven by reduced lean mass, not fat mass. Although Gipr expression was markedly reduced in visceral and subcutaneous adipose tissue from Gipradipo-/mice (41), Ap2/Fabp4 expression is not limited to adipocytes, because the Ap2/Fab4 promoter is transcriptionally active in heart, muscle, brain, macrophages, endothelium, and testis (46,51,52). We examined Gipr expression in adipose tissue depots and brain regions of independently generated Ap2-Cre:Gipr^{flx/flx} mice. Gipr transcripts were reduced in all adipose depots examined (Fig. 3A), as well as in the hypothalamus, hippocampus, and cortex (Fig. 3A). Ap2-Cre:Giprfix/fix mice weighed less at 8 weeks of age (Fig. 3B) and their weight remained below that of control mice throughout the HFD feeding period (Fig. 3C). However, the rate of weight gain between groups in response to HFD feeding was similar (Fig. 3D), as was body composition after 16 weeks of HFD feeding (Fig. 3E). Moreover, glucose tolerance (Fig. 3F), insulin tolerance (Fig. 3G), energy expenditure (Fig. 3H), food intake (Fig. 31), and tissue weights (Fig. 3J) were similar, although liver weight trended lower (P = 0.055). Thus, although reduction of Gipr in adipose tissue depots with Ap2-Cre is associated with modest changes in body weight, these findings cannot be directly attributed to reductions in adipose tissue Gipr expression, because the concurrent reduction in brain Gipr expression confounds attribution of phenotypes to adipose tissue in this mouse model.

Expression of *Gipr-Cre* Demonstrates Heterogeneity of Reporter Protein Expression Within Adipose Tissue

To reconcile our inability to reduce WAT *Gipr* expression with multiple *Adipoq-Cre* driver lines, we used *Gipr-Cre* to direct reporter protein expression. Crossing *Gipr-Cre* mice with a *Rosa26-LacZ* reporter line (53) produced abundant β -galactosidase activity in the pancreas, yet with little activity detected in WAT depots, including staining localized to blood vessels (Fig. 4). We next crossed *Gipr-Cre* mice with a *Rosa26-mT/mG* reporter line (54). Confocal microscopy of tissues from these mice demonstrated *Gipr* promoter activity in the pancreas, in line with islet GIPR expression (Fig. 4). *Gipr* promoter activity within WAT was much more heterogeneous, with a definitive signal seen in only a small fraction of putative adipocytes within multiple WAT depots.

Single-Nucleus RNA-Seq Localization of *Gipr/GIPR* Expression in Adipose Tissue

Collectively, the genetic findings in mice imply that Gipr expression within WAT is predominantly localized to nonadipocyte cell types. To further investigate this possibility, we analyzed Gipr/GIPR expression within distinct cell types of mouse and human inguinal and perigonadal adipose tissue using independently generated publicly available single nucleus RNA-seq data (34). Within mouse adipose tissue, Gipr mRNA was detected within Pdgfrbhigh pericytes, with minimal expression detected in Adipoq+ adipocytes (Fig. 5A-D). Glp1r was virtually absent in mouse adipose tissue (Fig. 5E); intriguingly, Glp2r was found in adipose stem and progenitor cells and macrophages (Fig. 5F). Gcgr showed an expression pattern similar to Gipr and was localized to pericytes (Fig. 5G). Similarly, scRNA-seq analysis of human subcutaneous and visceral adipose tissues detected GIPR primarily in *PDGFRB*^{high} pericytes, followed by *WT1*⁺ mesothelial cells, but not in $ADIPOQ^{high}$ adipocytes (Fig. 6A–E). Among the related class B, G protein-coupled receptors, only GLP2R, but not GLP1R and GCGR, was detected in some human adipose stem and progenitor cells and adipocytes (Fig. 6F-H). Hence, the available RNA-seq data are sufficiently useful for detection of GIPR and related class B, G protein-coupled receptor mRNA transcripts within various adipose tissue cell types, independently highlighting the lack of GIPR expression within the majority of mouse or human adipocytes.

DISCUSSION

Our current findings have implications for interpreting studies of GIP biology in adipose tissue. First, using independent mouse lines, we found that the murine *Gipr* is not expressed within the majority of adipocytes. Second, consistent with these findings, expression of Cre recombinase under control of the *Adiponectin* promoter does not meaningfully reduce *Gipr* expression in multiple adipose tissue depots. Third, interpretation of data generated using nonadipocyte-selective promoters to target adipose tissue *Gipr* expression, exemplified by *Ap2-Cre* (41), may be confounded by reduction of *Gipr* expression in multiple nonadipocyte cell types, including immune, neuronal, and endothelial cells (46). Although relative levels of *Gipr* mRNA transcripts were reported as normal in



Figure 3—*Ap2-Cre* expression results in *Gipr* knockdown in adipose and brain tissues. *A*) *Gipr* expression in adipose depots and brain tissues. (n = 4). *B*) Body weights (BWs) in 8-week-old mice prior to initiation of HFD feeding. (n = 7-11). Absolute (*C*) and percentage increases (*D*) in BWs of mice fed a 45% HFD diet for several weeks. (n = 7-11). *E*) Body composition expressed as a percentage of total (*left*) or absolute (*right*) weight in mice after 16 weeks of HFD feeding. (n = 8-11). *F*) Blood glucose levels and area under the glucose curve (AUC) during an oral glucose tolerance test (OGTT) after 17 weeks of HFD feeding. (n = 10-12). *G*) Blood glucose levels during an insulin tolerance test (ITT) after 6 weeks of HFD feeding. (n = 10-12). *H*) Oxygen consumption after 20 weeks of HFD feeding. (n = 8). *J*) Food intake (24 h) after 20 weeks of HFD feeding. (n = 8). *J*) Tissues weights after 20 weeks of HFD feeding. (n = 7-10). *A*) For relative mRNA expression, values were normalized to levels detected in RNA isolated from the same adipose tissue depots of *Ap2-Cre* mice. *P < 0.05 vs control.

the brain of $Gipr^{adipo-/-}$ mice (41), our analyses, using the same Ap2 promoter to express *Cre* and inactivate the *Gipr*, revealed substantial reduction of *Gipr* mRNA transcripts in

multiple regions of the murine central nervous system known to affect systemic metabolism. Taken together, these findings are consistent with a substantial proportion



Figure 4—Heterogeneous distribution of *Gipr* promoter activity in adipose depots. Representative whole-mount images of β -galactosidase activity (*left* column) and confocal fluorescence imaging (*right* column) of pancreas and adipose tissues from *Gipr Cre:Rosa26Lacz* and *Gipr Cre:mTmG* mice, respectively. GFP, Cre-positive green fluorescent protein; TdTomato, Cre-negative tomato red fluorescent protein.

of adipose tissue GIPR expression arising within nonadipocyte lineages.

In agreement with interpretation of the data obtained using genetic approaches in vivo, the single-nucleus RNAseq data provide further support for the concept that mouse and human adipocytes are not major sites of canonical *Gipr/ GIPR* expression. Indeed, pericytes appear to be a putative GIPR-expressing cell type in both human and mouse WAT, and additional human WAT *GIPR* expression is identified in mesothelial cells. These findings have implications for interpretation of the existing literature describing mechanisms of GIP action in adipose tissue and may generate new hypotheses about the actions of GIPR within WAT cell types that contribute to the biology of adipose tissue development and function.

The importance of understanding the biology of the adipose tissue GIPR and its impact has accelerated in part due to translational interest in targeting the GIPR for the treatment of obesity and diabetes (7). Indeed, GIPR agonism reduces food intake, body weight, and fat mass in HFD-fed mice through mechanisms requiring central nervous system GIPR activation (55). A GIP–GLP-1 co-agonist LY3298716, subsequently renamed tirzepatide, robustly stimulated cAMP accumulation in adipocyte-like cells derived from progenitors differentiated ex vivo, reduced food intake and adipose tissue mass, and produced substantial weight loss in both preclinical and clinical studies (56). Moreover, tirzepatide augmented adipose tissue glucose uptake and enhanced insulin sensitivity in a GLP-1R–independent manner in mice (57). Remarkably, GIPR blockade with antibodies directed against the mouse or human GIPR also reduced fat (WAT) mass, blocked the actions of exogenous GIP on human adipocytes ex vivo, and attenuated weight gain, without changes in lean mass in mice and nonhuman primates (27,31). Reconciliation of how both gain and loss of function at the GIPR produce overlapping effects on body weight, WAT mass, and function requires a more detailed understanding of how GIP controls metabolism and adipose tissue biology.

The results of several previous studies examining GIP action in adipose tissue have yielded conflicting results, with some studies demonstrating that GIP acts directly on WAT and other experiments invoking a role for GIP as an insulin sensitizer on adipocytes, using cells differentiated from adipocyte progenitors ex vivo (11,58,59). Indeed, the very slow kinetics of the adipose tissue response to GIP (59) have prompted the suggestion that GIP might act indirectly on adipocyte lipid metabolism, through one or more downstream mediators such as insulin or resistin (35,60,61). Nevertheless, substantial data suggest that adipocyte-like cells studied ex vivo express a functional GIPR coupled to cAMP accumulation and fatty acid uptake (43,57).



Figure 5—Mouse single-nucleus RNA-seq data were used to localize WAT *Gipr* expression to pericytes. *A*) Single-cell nucleus RNA-seq data (34) from a range of mouse adipose tissue cell types were analyzed for expression of *B*) *Gipr*; *C*) *Adipoq*; *D*) *Pdgfrb*, a gene expressed in endothelial/pericyte and adipose tissue progenitors; *E*) *Glp1r*; *F*) *Glp2r*; and *G*) *Gcgr*. ASPC, adipocyte stem progenitor cell.

Previous studies using mouse genetics to interrogate the role of the adipocyte GIPR have been partially inconclusive. For example, transgenic targeting of GIPR expression to WAT of $Gipr^{-/-}$ mice using the Ap2/Fabp4promoter produced weight gain independent of changes in fat mass, without any meaningfully evident metabolic phenotypes (62). Conversely, reduction of WAT GiprmRNA transcripts using the Ap2/Fabp4 promoter to direct *Cre* expression to several cell types, including adipocytes, reduced WAT *Gipr* expression in mice with lower body weight and lean body mass, yet without change in fat mass (41). More recent studies using *Adipoq-Cre* to target the mouse adipocyte *Gipr* revealed loss of *Gipr* expression in adipocytes differentiated ex vivo, together with reduced GIP-stimulated cAMP accumulation and decreased fatty acid uptake (43). Notably, however, the levels of *Gipr* mRNA within WAT depots from *Gipr*^{Adipo-/-} mice were not reported.



Figure 6—Human single-nucleus RNA-seq data were used to localize WAT *GIPR* expression to pericytes and mesothelial cells. *A*) Singlecell nucleus RNA-seq data (34) from a range of human adipose tissue cell types were analyzed for expression of *B*) *GIPR*; *C*) *ADIPOQ*; *D*) *PDGFRB*; *E*) *WT1*, a marker for the mesothelial cell lineage; *F*) *GLP1R*; *G*) *GLP2R*; and *H*) *GCGR*. ASPC, adipocyte stem progenitor cell.

The growing importance in understanding the actions of GIP in WAT is further augmented by interest in the mechanisms of action of tirzepatide (63). Administration of tirzepatide or a long-acting GIPR agonist improved insulin sensitivity, associated with enhanced glucose uptake into WAT (57). Interestingly, however, RNA-seq analysis of WAT from mice treated with tirzepatide or a long-acting GIPR agonist showed no changes in metabolic gene expression within WAT depots, whereas a GIPR agonist and tirzepatide differentially regulated >1,000 genes within BAT (57). These latter findings are consistent with a functional role for the canonical murine GIPR in regulation of genes important for thermogenesis, lipid metabolism, and cytokine expression in BAT (47,61).

Limitations and Future Perspectives

Our data require interpretation with caution due to a number of important limitations. First, we focused almost entirely on mRNA expression, because of the lack of suitably validated antisera for detection of the mouse GIPR (32,33). Data from experiments using reporter genes to infer expression should be considered with caveats, because the readouts may reflect activation of transcriptional sequences in one or more early adipose tissue lineages that subsequently give rise to differentiated adipocytes. Hence, whether reporter gene expression within a few adipocytes coincides with simultaneous coexpression of the Gipr mRNA transcript or protein in the same differentiated cell remains uncertain. Although the scRNA-seq and gene-targeting data align with the concept that mouse adipocyte *Gipr* expression is uncommon, we did not study adipose tissue depots from a wide range of mice with metabolic perturbations. For example, animals with diabetes, insulin deficiency or resistance, or obesity, might exhibit upregulation of adipocyte Gipr expression—scenarios that require additional investigation.

It is also worth noting that low-level adipocyte expression of class B GPCRs such as *Glp1r* or *Gipr* might not easily be detected using thresholds set for scRNA-seq, although the same analyses successfully detected Gipr and Glp2r mRNAs in nonadipocyte cell types within adipose tissue (34). Finally, our data do not rule out an important role for GIPR activity in adipose tissue. The small fraction of mature GIPR⁺ adipocytes identified by scRNA-seq could represent a key subset of cells that contribute to regulation of overall adipose tissue function through paracrine or endocrine processes. Alternatively, the colocalization of Gipr/GIPR with Pdgfrb/PDGFRB populations may represent early precursor cells, potentially suggesting a role for GIP in preadipocyte function. Finally, there is much less information available on the cellular localization of GIPR in human adipose tissue depots across the life span in people living with diabetes or obesity, hence the putative importance of adipocyte expression of the human GIPR requires greater scrutiny. In summary, our data introduce further complexity in conceptualizing how gain or loss of GIPR signaling affects adipose tissue biology and adipocytes in vivo.

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