GIPR agonism mediates weight-independent insulin sensitization by tirzepatide in obese mice

Ricardo J. Samms, … , Kyle W. Sloop, William C. Roell


Tirzepatide (LY3298176), a dual GIP and GLP-1 receptor (GLP-1R) agonist, delivered superior glycemic control and weight loss compared with GLP-1R agonism in patients with type 2 diabetes. However, the mechanism by which tirzepatide improves efficacy and how GIP receptor (GIPR) agonism contributes is not fully understood. Here, we show that tirzepatide is an effective insulin sensitizer, improving insulin sensitivity in obese mice to a greater extent than GLP-1R agonism. To determine whether GIPR agonism contributes, we compared the effect of tirzepatide in obese WT and Glp-1r–null mice. In the absence of GLP-1R–induced weight loss, tirzepatide improved insulin sensitivity by enhancing glucose disposal in white adipose tissue (WAT). In support of this, a long-acting GIPR agonist (LAGIPRA) was found to enhance insulin sensitivity by augmenting glucose disposal in WAT. Interestingly, the effect of tirzepatide and LAGIPRA on insulin sensitivity was associated with reduced branched-chain amino acids (BCAAs) and ketoacids in the circulation. Insulin sensitization was associated with upregulation of genes associated with the catabolism of glucose, lipid, and BCAAs in brown adipose tissue. Together, our studies show that tirzepatide improved insulin sensitivity in a weight-dependent and -independent manner. These results highlight how GIPR agonism contributes to the therapeutic profile of dual-receptor agonism, offering mechanistic insights into the clinical efficacy of tirzepatide.

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Introduction

Type 2 diabetes mellitus (T2DM) is a global pandemic, placing significant strain on healthcare systems world wide (1). Driven primarily by excess body weight, systemic insulin resistance (IR) is a key driver of pancreatic β cell dysfunction and the subsequent progression to T2DM (2, 3). Although the etiology remains to be confirmed, current evidence implicates several factors (e.g., genetic predisposition, hyperlipidemia/lipotoxicity, impaired branched-chain amino acid (BCAA) catabolism, inflammation, endoplasmic reticulum stress, and mitochondrial dysfunction) in the development of whole-body IR (4). To date, caloric restriction–mediated weight loss, bariatric surgery, or pharmacotherapy with thiazolidinediones (TZDs, small-molecule agonists of the nuclear hormone receptor PPARγ) are the only effective approaches that specifically treat IR for improving glucose control (5). However, in the long term, caloric restriction is ineffective due to counterregulatory responses and compliance issues, bariatric surgery is an expensive procedure carrying significant risk, and the TZD drug class has fallen out of favor due to an unacceptable side effect profile (e.g., weight gain, edema, fracture, ref. 6). Therefore, new therapeutic agents that are capable of targeting multiple organs/signaling pathways to enhance insulin sensitivity may offer an attractive mechanism-based approach for treating T2DM (7).

Over the past decade, glucagon-like peptide-1 receptor agonists (GLP-1RAs) have emerged as popular medicines for the treatment of T2DM due to their safety and efficacy (8). The glycemic benefits associated with GLP-1RAs are due to enhanced β cell function, delayed gastric emptying, and weight loss–driven improvements in insulin sensitivity (9). However, although the GLP-1RA class has improved the standard of care, many patients still do not reach their glycemic targets (10), and thus, there remains a need to identify agents capable of complementing the actions of GLP-1RAs. As key sites of energy storage and oxidation, white and brown adipose tissue (WAT and BAT) are at the forefront of the link between obesity-induced IR and the subsequent development of T2DM (11, 12). Thus, one strategy to augment the therapeutic profile of GLP-1RAs is to complement their actions with agents that target WAT and BAT to improve their capacities for metabolic substrate disposal, thus lowering plasma levels of lipids, BCAAs, and other metabolites that may contribute to systemic IR (13, 14). Glucose-dependent insulinotropic polypeptide (GIP) is a hormone released from the gut following the consumption of food (15). Similar to GLP-1, the primary physiological role of GIP is to enhance glucose-dependent insulin secretion (16, 17). However, the metabolic effects of GIP extend beyond the pancreas and include several metabolically relevant organs (18). In contrast to the glucagon-like peptide 1 receptor

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benefits associated with chronic GIPR engagement are linked to improved adipose tissue health, characterized by enhanced adipocyte insulin signaling, reduced proinflammatory immune cell infiltration, recruitment of metabolic pathways associated with lipid storage and oxidation, and the release of insulin-sensitizing adipokines (23–26). Together, the ability of GIP to target (GLP-1R), the GIP receptor (GIPR) is expressed in WAT (19) and BAT (20). Via its effects in these tissues, GIP has been described primarily as a regulator of glucose and lipid metabolism (20–22). Overexpression of GIP or administration of long-acting GIPR agonists (LAGIPRAs) enhances whole-body insulin sensitivity in models of obesity-induced IR (23, 24). Mechanistically, the benefits associated with chronic GIPR engagement are linked to improved adipose tissue health, characterized by enhanced adipocyte insulin signaling, reduced proinflammatory immune cell infiltration, recruitment of metabolic pathways associated with lipid storage and oxidation, and the release of insulin-sensitizing adipokines (23–26). Together, the ability of GIP to target

![Figure 1. Chronic treatment with tirzepatide enhanced insulin tolerance in obese mice.](image-url) Obese insulin-resistant mice were dosed once daily for 14 days with vehicle or tirzepatide (TZP, 10 nmol/kg; n = 5–8 per group). (A) Daily body weight and food intake. (B) Tissue weights, (C) plasma, (D) leptin triacylglycerides and free fatty acids (FFA), and (E) liver triglyceride after 14 days of treatment. (F) Fed and (G) fasted blood glucose and plasma insulin. (H) Fed adiponectin and insulin-like growth factor binding protein 2 (IGFBP2). (I) Insulin tolerance test. Data are presented as mean ± SEM. *P < 0.05 compared with vehicle. Statistical analyses performed included (A) 2-way ANOVA, (B–H) Student’s unpaired t test, and (I) 1-way ANOVA followed by Dunnett’s multiple comparisons test, where appropriate.
adipose tissue and improve metabolic homeostasis highlights its attractiveness for partnering with GLP-1 (18, 22). As a therapeutic modality, next-generation antidiabetic agents engineered to simultaneously target the GIP and GLP-1 receptors (GLP-1Rs) are being pursued (27–29). The most advanced multireceptor agonist is tirzepatide (LY3298176), an imbalanced and biased GIP and GLP-1R agonist (28, 30). In a phase 2b trial of patients with T2DM, tirzepatide delivered superior weight loss and improved glycemic control when compared with a selective GLP-1RA ( dulaglutide, 1.5 mg; ref. 31). Importantly, improved glucose homeostasis was associated with enhanced insulin sensitivity, and post hoc analysis of this study suggested that the majority of the insulin sensitization occurred independent of weight loss (32). Therefore, we investigated the mechanistic basis of how tirzepatide may improve insulin sensitivity and whether GIPR agonism directly contributes to this benefit.

Results
To determine if tirzepatide improves insulin sensitivity, we conducted a series of chronic studies in high-fat diet–fed obese IR mice. Treatment with tirzepatide reduced body weight, food intake, tissue weights, and plasma leptin (Figure 1, A–C). Fourteen days of treatment with tirzepatide reduced circulating triglyceride levels and free fatty acids (FFAs) and lowered hepatic lipid content (Figure 1, D and E). In line with improved glucose homeostasis, tirzepatide reduced fed (Figure 1F) and fasted (Figure 1G) blood glucose and plasma insulin, while increasing circulating levels of the insulin sensitizers adiponectin and IGFBP2 (refs. 33, 34 and Figure 1H). Furthermore, in accordance with a state of heightened insulin sensitivity, tirzepatide improved insulin tolerance (Figure 1I). Importantly, chronic treatment with tirzepatide reduced body weight and food intake and enhanced markers of insulin sensitivity when dosed at room temperature (24°C, Figure 1) and ther-
Glucose uptake. Tirzepatide increased insulin-stimulated glucose disposal in skeletal muscle (Figure 2, C–E) and WAT (Figure 2, F and G). These experiments indicate that tirzepatide is a highly effective insulin sensitizer in obese IR mice, with enhanced insulin-stimulated glucose disposal in skeletal muscle and adipose tissue, appearing to account for the glycemic benefits.

One of the most effective approaches to achieving clinically meaningful improvements in glycemic control is weight loss (35). Therefore, to determine if the effect of tirzepatide on systemic insulin sensitivity is driven in a weight-dependent and/or -independent manner, we compared the insulin-sensitizing action of tirzepatide to a selective GLP-1RA (semaglutide), a mechanism that has been shown to ameliorate IR entirely due to weight loss.
Figure 4. GIPR agonism contributes to the weight-independent insulin sensitization action of tirzepatide. WT (C57BL/6J) and germline, whole-body, glucagon-like peptide 1 receptor–KO mice (Glp-1r−/− mice) were individually housed in a temperature-controlled (27°C) environment with a 12-hour light/12-hour dark cycle and fed a high-fat (60% of calories from fat) diet for 12 weeks. (A) Weekly body weight and food intake. (B) Fat and lean mass after 12 weeks of high-fat feeding. Obese insulin-resistant Glp-1r−/− mice, dosed once daily with either vehicle (n = 12–15) or tirzepatide (TZP, 10 nmol/kg) for 14 days. (C) Daily body weight and food intake. (D) Fasting blood glucose and plasma insulin following 14 days of treatment. Following 14 days of treatment, insulin sensitivity was assessed via a hyperinsulinemic-euglycemic clamp. (E) Average glucose infusion rate (GIR) and that during the final 30 minutes of the clamp. (F) Endogenous glucose production (EGP). Insulin-stimulated glucose disposal in (G) soleus, (H) red, and (I) white gastrocnemius skeletal muscle as well as (J) epididymal white adipose tissue (eWAT) and (K) inguinal white adipose tissue (iWAT). Data are presented as mean ± SEM. *P < 0.05 compared with vehicle. Statistical analyses performed included Student’s unpaired t test, 2-way ANOVA, where appropriate.
(36). For these studies, we matched weight loss in obese IR mice by dosing tirzepatide (3 nmol/kg) at a 3-fold lower dose than semaglutide (10 nmol/kg). To further control for weight-driven improvements in insulin sensitivity, we pair fed (PF) a group of animals the same daily food intake as animals dosed with tirzepatide. Chronic administration of either agent reduced body weight and food intake in obese IR mice (Figure 3A). Following 14 days of treatment, there was no difference in weight loss in animals treated with tirzepatide versus those treated with semaglutide. Tirzepatide reduced fasting glucose levels lower than those seen in semaglutide-treated or PF animals (Supplemental Figure 1G). Tirzepatide, semaglutide, and PF reduced fasting insulin when compared with that in vehicle-treated animals (Supplemental Figure 1H). To compare the insulin-sensitizing effects of the 2 treatments, we conducted hyperinsulinemic-euglycemic clamp experiments following the 14 days of dosing. Strikingly, despite eliciting equivalent weight loss, administration of tirzepatide increased whole-body insulin sensitivity more than that achieved in the semaglutide treatment and PF groups (Figure 3B). Tirzepatide increased the GIR by 4.7-fold, while the semaglutide and PF groups increased the GIR by 3.5-fold and 3-fold, respectively, when compared with the vehicle-treated group. There was a 1.4-fold and 1.5-fold increase in the GIR in animals administered tirzepatide when compared with treatment with semaglutide or PF, respectively (Figure 3C). Importantly, comparing the insulin-sensitizing action of tirzepatide to that of the weight-matched groups (semaglutide and PF) indicated that the weight-independent effect of tirzepatide accounts for 30% of the total improvement in insulin sensitivity in obese IR mice. No effect on EGP was observed for any of the groups (Figure 3D). Tirzepatide augmented insulin-stimulated glucose disposal in skeletal muscle.
Table 1. In vitro potency and pharmacokinetic analysis

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<th>mGLP-1R (nM)</th>
<th>mGIPR (nM)</th>
<th>mGCGR (nM)</th>
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<tr>
<td></td>
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<td>Dose (nmol/kg)</td>
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<td>Cmax (nmol/L)</td>
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<tr>
<td>Tmax (h)</td>
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<td>AUC∞ (h nmol/L)</td>
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<td>CL (i.v.) or CL/F (s.c.) (mL/h/kg)</td>
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</tr>
<tr>
<td>t1/2 (h)</td>
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<td>6.29</td>
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In vitro potency and pharmacokinetic analysis of a long-acting glucose-dependent insulinotropic polypeptide receptor agonist (LAGIPRA) in CD-1 mice. In vitro potency values for LAGIPRA in HEK293 cells expressing mouse GIPR (mGIPR), glucagon-like peptide 1 (mGLP-1R), or glucagon (mGCGR) receptor. Data are presented as mean ± SEM. AUC∞, area under the curve from 0 to infinity; CL, clearance; CL/F, apparent clearance; Cmax, maximal concentration; Tmax, time at maximal concentration; t1/2, elimination half-life.

To elucidate the potential mechanism(s) by which tirzepatide (and GIPR agonism) improved insulin sensitivity, we conducted RNA-Seq analysis in several tissues. Interestingly, there was no effect of tirzepatide or LAGIPRA on metabolic genes in skeletal muscle or WAT (data not shown). Unexpectedly, the most marked differences were found in BAT, where both tirzepatide and LAGIPRA treatment robustly induced the expression of genes associated with the catab-
and/or T2DM (41–43). Here, a total of 3671 and 1908 genes were found to be differentially regulated by treatment with tirzepatide or LAGIPRA, respectively, in BAT (Figure 7A). Overall, 1926 and 1296 genes were upregulated and 1745 and 612 genes were downregulated by tirzepatide and LAGIPRA, respectively. Consistent with GIPR

**Figure 6. LAGIPRA improves insulin sensitivity in obese insulin-resistant mice.** High-fat diet–fed obese insulin-resistant mice dosed daily with either vehicle (n = 8–14) or a long-acting glucose-dependent insulinotropic polypeptide receptor agonist (LAGIPRA, n = 8–14) for 14 days. (A) Daily body weight and food intake. (B) Fasting blood glucose and plasma insulin following 14 days of treatment. Hyperinsulinemic-euglycemic clamp following 14 days of treatment. (C) Average glucose infusion rates throughout and during the final 30 minutes of clamp (GIR). (D) Endogenous glucose production (EGP). Insulin-stimulated glucose disposal in (E) soleus, (F) red, and (G) white gastrocnemius skeletal muscle and (H) epididymal white adipose tissue (eWAT) and (I) inguinal white adipose tissue (iWAT). Data are presented as mean ± SEM. *P < 0.05 compared with vehicle. Statistical analyses performed included Student’s unpaired t test, 2-way ANOVA, or Kruskal-Wallis test, where appropriate.
engagement, 703 upregulated and 198 downregulated genes were shared between tirzepatide and LAGIPRA treatment. Accordingly, gene set enrichment analysis of differentially expressed genes indicated that tirzepatide and LAGIPRA affected similar molecular targets, including pathways associated with cellular and mitochondrial metabolism. Specifically, tirzepatide and/or LAGIPRA treatment induced the mRNA expression of metabolic transcriptional regulators (ChREBP, SREBP-1, RXRβ, PPARγ, and KLF-15; Figure 7B and Supplemental Figure 3) and recruited genes associated with the uptake, storage, breakdown, and oxidation of glucose and lipids in BAT (GLUT1, GLUT4, ATGL, DGAT1, and DGAT2; Supplemental Figure 3). Interestingly, tirzepatide and LAGIPRA treatment induced the expression of BCAT2 and BCKDH, enzymes that catalyze the first and second steps of BCAA catabolism, respectively (Figure 7B). Furthermore, LAGIPRA treatment, in particular, induced expression of genes associated with downstream steps in BCAA catabolism (e.g., HIBCH, HIBADH, ACAD8, ACADM, IVDC, MCEE, and MCCC2; Figure 7B). Consistent with increased TCA cycle flux and electron transport, tirzepatide and/or LAGIPRA treatment stimulated the recruitment of genes associated with mitochondrial substrate transport (e.g., CPT1A, MIPC2, MCAT, SLCS2A39, SLCS2A11, SLCS2A22, and SLCS2A29), fatty acid oxidation (PDK2, ACAD8, ACAD9, ACAD10, ACADL, ACADVL, and ACADM), electron transport activity (COX5A, COX5B, COX7A1, COX8b, COX17, COX6B1, COX10, and COX11), and thermogenic capacity (UCP1; Supplemental Figure 3). There was no effect of PF on several of these genes, suggesting that GIPR agonism mediates a direct and weight-independent induction of metabolic pathways associated with the oxidation of glucose, lipids, and BCAAs in BAT.

Of particular interest, we found that TZP modulated BCAA metabolism in BAT. To determine if the induction of BCAA catabolic genes in BAT resulted in a functional outcome, we investigated the effect of tirzepatide and LAGIPRA on tissue and circulating levels of BCAAs. There was an increase in BCAAs (valine, leucine, and isoleucine) in BAT of mice treated with either tirzepatide or LAGIPRA (Table 2). Furthermore, indicative of tissue level catabolism of BCAAs, there was an increase in the BCAA catalytic product glutamate (a product of the first step of BCAA catabolism) in BAT of mice treated with tirzepatide (Table 2). In line with tissue changes affecting circulating BCAA, tirzepatide treatment decreased plasma levels of BCAAs (valine, leucine, and isoleucine) and BCAA catalytic products glutamate and branched-chain α-ketoacids (BCKAs; α-ketoisocaproate and α-keto-β-methylvalerate; Table 2). Interestingly, while there was no effect of LAGIPRA treatment on BCAA levels, but consistent with the robust effect of LAGIPRA on BCAA catabolic genes, LAGIPRA decreased circulating levels of BCKAs (α-ketoisocaproate and α-keto-β-methylvalerate; Table 2). The translational relevance of these preclinical studies is exemplified by post hoc analysis of clinical data, suggesting that the weight-independent insulin-sensitizing action of tirzepatide is strongly associated with reduced circulating BCAA/BCKA in patients with T2DM (44).
Glp-1r–/– mice. Using ity in a weight-dependent and -independent manner in obese IR whether the GIPR contributes to the efficacy. The major findings sensitivity beyond what occurs by weight loss alone and, importantly, ty-induced IR to investigate how tirzepatide improves insulin sen-
tissue-specific, and molecular levels in a mouse model of obesi-
sive approach, combining measurements at the whole-body, metabolic pathways associated with the oxidation of glucose, lip-
id, and BCAAs in adipose tissue.

Discussion

Multireceptor pharmacology is at the forefront of next-genera-
tive therapies for the treatment of metabolic diseases (27, 45). One of these investigational agents is tirzepatide, a dual GIP and GLP-1R agonist, which has shown enhanced glycemic control and weight loss in patients with T2DM when compared with GLP-1R agonist monotherapy therapy (31). Here, we took a comprehen-
approach, combining measurements at the whole-body, tissue-specific, and molecular levels in a mouse model of obesity-induced IR to investigate how tirzepatide improves insulin sensitivity beyond what occurs by weight loss alone and, importantly, whether the GIPR contributes to the efficacy. The major findings of the current report are that tirzepatide improves insulin sensitivity in a weight-dependent and -independent manner in obese IR mice. Using Glp-1r–/– mice, we showed that GIPR agonism accounts for the weight-independent insulin sensitization. Further, mecha-
nistic studies indicate that tirzepatide-mediated improvements in insulin sensitivity are associated with the induction of glucose, FFA, and BCAA oxidation in BAT.

Obesity-induced IR is a key component underlying the develop-
ment and progression of T2DM (46). Currently, there are limited efficacious, safe, and durable medications that target this dysfunction in T2DM (47). One of the most effective therapies for the treatment of T2DM is weight loss (48, 49). In line with this, GLP-1R agonists drive weight loss and improve insulin sensitivity (36). However, a caveat of improving insulin sensitivity principally through weight loss is that once body weight is regained, metabolic benefits may be compromised (50, 51). Thus, improving insulin action in both a weight-dependent and -independent manner could provide greater treatment efficacy and durability. A key finding of the current study is that tirzepatide ameliorated peripheral IR in a model of DIO. Importantly, the insulin sensitization was underscored by enhanced glucose disposal in skeletal muscle and in both visceral and s.c. WAT. Consequently, a key question arising from this study was whether a component of the observed insulin sensitization occurred independent of weight loss. To address this, we matched the body weight of animals treated with tirzepatide to that of a PF group and with animals dosed with a selective GLP-1R monoagonist (semaglutide), a mechanism that improves insulin action primarily due to weight loss (36). Remarkably, despite losing equivalent body weight, animals treated with tirzepatide had a greater improvement in systemic insulin sensitization when compared with the weight-matched groups. Moreover, this weight-independent insulin benefit was coupled with augmented glucose disposal in skeletal muscle and in both WAT and BAT. Collectively, these results suggest that tirzepatide delivers both weight-dependent and -independent insulin sensitization in obese IR mice. The clinical implications of these findings are that the weight-independent insulin sensitization identified here could offer greater treatment durability, in contrast to agents that provide only weight-driven benefits. In support of this hypothesis, HOMA-IR analysis has suggested that a significant proportion of tirzepatide-induced insulin sensitization occurs in the absence of weight loss in subjects with T2DM (52). Together with studies showing that GIP can improve insulin sensitivity (23, 24), this prompted the hypothesis that engagement of the GIPR may drive the weight-independent insulin sensitization action of tirzepatide.

Two therapeutic modalities have been proposed to target the GIPR for treating metabolic diseases (22, 53). Originally, the discovery that Gipr–/– mice are protected from obesity (54), in addition to studies showing that GIP enhances lipid storage in WAT (55, 56), led to the idea that GIP was obesogenic and consequently prompt-
ed several approaches to block GIP action (57). Indeed, antagoniz-
ing antibodies targeting the GIPR have been shown to protect mice from DIO (58, 59). To date, however, these agents have not been shown to drive weight loss in already obese animals; further, GIPR antagonism offers minimal glycemic benefit (53, 60). In addition, as GIP is the primary incretin in humans (17, 61), blockade of GIP action may be detrimental to glycemic control, while chronic GIPR agonism either has no effect on body weight or drives weight loss and improves insulin action in mice (23, 24, 28, 29, 62, 63). Here, to determine if GIPR agonism accounts for the weight-independent insulin sensitization, we treated obese IR Glp-1r–/– mice chronically

<table>
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<th>Metabolite</th>
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<th>Plasma</th>
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<tr>
<td></td>
<td>TZP (3 nmol/kg)</td>
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Tirzepatide and LAGIPRA reduced circulating BCAA in obese IR mice. High-fat diet–fed obese insulin-resistant mice (C57BL/6J) were dosed once daily with vehicle (saline, n = 5–6), tirzepatide (TZP, n = 5–6), or a long-acting glucose-dependent insulino
tropic polypeptide receptor agonist (LAGIPRA, n = 5–6). Following 14 days of treatment, plasma and brown adipose tissue samples were collected for plasma and tissue branched-chain amino acid (BCAA) analysis. Data are presented as fold change compared with vehicle. *n = 0.05 compared with vehicle. Statistical analyses were performed using a Student’s unpaired t test.

Taken together, these data indicate that tirzepatide (and GIPR agonism) may improve systemic insulin sensitivity by activating metabolic pathways associated with the oxidation of glucose, lip-

ids, and BCAAs in adipose tissue.
with tirzepatide, and to complement this strategy, we dosed obese IR WT animals with a LAGIPRA. Consistent with data from other reports (23, 28, 29, 62), neither LAGIPRA nor tirzepatide (in GLP-1R-KO animals) reduced body weight, supporting the hypothesis that GLP-1R agonism is required to unlock the synergistic action GIPR agonism on weight loss (18). Importantly, this provides an experimental paradigm to investigate the weight- and GLP-1R– independent effect of tirzepatide (GIPR agonism) on systemic insulin sensitivity. Using hyperinsulinemic-euglycemic clamp studies, we found that both LAGIPRA and tirzepatide improved insulin sensitivity in obese mice in the absence of weight loss. Notably, the magnitude of enhanced insulin action matched the weight-independent insulin sensitization occurring when comparing tirzepatide to semaglutide. Further, in line with the GIPR being expressed in adipose tissue (20), we found that for both LAGIPRA and tirzepatide (in Glp-1r–/– mice) enhanced insulin sensitization was underlined by augmented glucose disposal in WAT. Thus, although the specific contribution of WAT to total postprandial glucose clearance is not comparable to that of skeletal muscle (64, 65), the cumulative findings indicate that treatment with tirzepatide results in weight-independent insulin sensitivity in obese IR mice and that engagement of the GIPR accounts for this benefit. These data agree with previous findings showing that GIPR agonism augments insulin sensitivity in obese mice (24). Mechanistically, GIP has been shown to improve insulin sensitivity due to the recruitment of metabolic pathways associated with the storage and oxidation of glucose and FFAs in adipose tissue (23, 24).

Although the cause of obesity-associated IR is intensely debated (4, 66, 67), recently, a strong association between elevated BCAAs/BCKAs, obesity, systemic IR, and T2DM has consistently been reported (66, 68). This has led to the suggestion that dysregulation of BCAA metabolism may play a causal role in developing IR (66). Indeed, dysregulation of BCAA catabolism in adipose tissue plays a role in the rise of circulating BCAAs due to reductions in the expression of enzymes in BCAA catabolism (69–71). Consequently, in models of obesity-induced IR, induction of BCAA catabolism either systemically or specifically in adipose tissue, reduces circulating BCAAs and alleviates IR (39, 67–69). In accordance, we showed that tirzepatide-mediated (and GIPR agonism–mediated) improvements in insulin sensitivity are coupled with the induction of enzymes responsible for the catabolism of BCAAs in BAT. Specifically, both tirzepatide and LAGIPRA were found to increase expression of genes linked with glucose and lipid metabolism. The most robust effect occurred on the expression of BCAT2 and BCKDH, enzymes that catalyze the first and second (and rate limiting) steps in BCAA catabolism, respectively, as well as several downstream members of the BCAA catabolic pathway in BAT. Importantly, these molecular changes were found to be associated with increased tissue levels of BCAAs and reduced circulating levels of BCAAs and their product BCKA. Our studies are supported by data showing the GIPR is expressed in BAT and that GIP can act directly on brown adipocytes to induce metabolic gene expression (20). In addition, the potential clinical relevance of our findings is highlighted by post hoc analyses showing that tirzepatide reduces circulating BCAAs in patients with T2DM and that this effect is strongly associated with biomarkers indicative of enhanced insulin sensitivity (44). These results lead to the intriguing hypothesis that one way in which tirzepatide may enhance insulin sensitivity in patients is through the catabolism of glucose, lipids, and BCAAs in BAT. In support of this, activated BAT is reported to oxidize circulating glucose, lipids, and BCAAs in BAT in humans (39). Therefore, while BAT has classically been considered to have a limited effect on human metabolism (72, 73), studies show that adult humans have functional BAT (74), which when activated, is capable of positively affecting systemic metabolism (43, 75, 76). This clearly highlights the pharmacological potential of this organ in humans (77) and is in line with recent clinical findings indicating that the presence of BAT in humans is associated with cardiometabolic health (78). Thus, while it is likely to make a smaller contribution in humans (compared with mice), our data suggest that BAT may aid weight-dependent mechanisms in the lowering of circulating lipids and BCAAs, enhancing the improvement of insulin sensitivity by tirzepatide in patients with T2DM (44, 52). We note that mechanistic studies in human subjects are further required to fully test this hypothesis.

Summary. In summary, here we demonstrate that tirzepatide and LAGIPRA improve insulin sensitivity in obese IR mice. Importantly, we found that tirzepatide enhances insulin action to a greater extent than GLP-1RA treatment and that this additional efficacy occurs independent of changes in body weight. Using two complementary approaches, we demonstrate that this weight-independent insulin sensitization is accounted for by engagement of the GIPR. Mechanistically, tirzepatide appears to mediate its insulin-sensitizing action in a weight-dependent mode of action (via GIPR agonism) manner by the induction of metabolic pathways linked to the oxidation of glucose, lipids, and BCAAs. Together, this reduces excess nutrient delivery to metabolically relevant organs, subsequently enhancing systemic insulin sensitivity. The weight-independent insulin sensitization identified here could offer greater treatment durability, in contrast to agents that provide only weight-driven effects. Further, these preclinical data demonstrate how GIPR agonism contributes to enhanced glycemic control of tirzepatide, a dual GIP and GLP-1R agonist.

Methods

Peptide synthesis and in vitro characterization. LAGIPRA, GLP-1R agonist semaglutide, and tirzepatide were synthesized at Eli Lilly and Company. cAMP assays were conducted in HEK293 cells expressing the mouse GLP-1R, GIPR, or glucagon receptor (GcgR). Using homogeneous time-resolved fluorescence methods to assess cAMP accumulation (CisBio CAMP Dynamic HTRF Assay Kit, 62AM4PEJ), assays were conducted to determine the potency of LAGIPRA performed in the presence of cascin (instead of serum albumin) as a nonspecific blocker, which does not interact with the fatty acid moiety of LAGIPRA. Intracellular cAMP levels were determined by extrapolation using a standard curve. Dose-response curves of compounds were plotted as the percentage of stimulation normalized to minimum (buffer only) and maximum (maximum concentration of each control ligand) values and analyzed using a 4-parameter nonlinear regression fit with a variable slope (Genedata Screener 13). EC50 is the concentration of compound causing half-maximal stimulation in a dose-response curve.

Animals. All animals were individually housed in a temperature-controlled (25°C ± 0.9°C to 27°C ± 0.9°C) facility with a 12-hour-light/12-hour-dark cycle. WT, Glp-1r–/–, and Gipr–/– mice on a C57BL/6 genetic background (28, 79) were maintained at Taconic Contract Breeding.
High-fat diet study in WT and Glp-1r−/− mice. WT and Glp-1r−/− mice housed in a thermal neutral environment (27°C) were fed a 60% high-fat diet (D12492; Research diets) for a minimum of 12 weeks. Body composition was assessed before and after high-fat feeding using quantitative nuclear magnetic resonance. Body weight and food intake were recorded weekly.

Chronic dosing studies in obese insulin-resistant WT animals. Male mice were maintained on a high-fat diet (consisting of 60% fat, 20% carbohydrate, and 20% protein caloric content D12492; Research Diets) and had free access to food and water before randomization by weight. Animals received s.c. injections of vehicle (n = 6–16), semaglutide (10 nmol/kg, n = 16), or tirzepatide (3 and 10 nmol/kg, n = 6–16) or LAGIPRA (300 nmol/kg) once a day for 14 days. Mice were sacrificed, plasma was collected in EDTA-coated tubes, and tissues were collected and snap frozen in liquid nitrogen.

Chronic treatment with tirzepatide in obese insulin-resistant Glp-1r−/− mice. Male Glp-1r−/− mice were fed a high-fat diet (60% of calories from fat, D12492; Research diets) for a minimum of 16 weeks. Following this period animals were weighed and body composition was determined using quantitative nuclear magnetic resonance. Mice were then randomly assigned to treatment groups in which they received a daily injection of vehicle (n = 16) or tirzepatide (10 nmol/kg, n = 16) for 14 days. Body weight and food intake were recorded daily. Fourteen days after the start of each experiment, insulin sensitivity was determined via insulin clamp.

In vivo metabolic analysis. Metabolic rate and substrate utilization rates were determined in obese mice treated with vehicle (n = 6) or LAGIPRA (300 nmol/kg, n = 6) using an open respirometer system (LabMaster System; TSE Systems). Briefly, oxygen consumption (VO₂, mL/kg/h) and carbon dioxide production (VCO₂, mL/kg/h) were measured throughout the final 7 days of a 14-day dosing period. VO₂ (mL/kg/h) and VCO₂ (mL/kg/h) were used to calculate energy expenditure and respiratory exchange ratio (RER = VCO₂/VO₂). Animals had ad libitum access to water and high-fat diet when in metabolic cages. All measurements were made at approximately 24°C to 25°C.

Insulin clamp. Catheters were placed in the left carotid artery and the right jugular vein 7–8 days prior to euglycemic clamp. Animals were fasted overnight and, after 2 hours of study box acclimation, a bolus/continuous infusion of 3-H-glucose (PerkinElmer; 6 μCi bolus; basal period, 0.05 μCi/min; clamp period, 0.125 μCi/min) was initiated and maintained throughout the test period. Blood glucose was measured from arterial blood via a glucometer every 5 minutes during the clamp period. An i.v. infusion of 3 mIU/kg/min Humulin R (Eli Lilly and Company) and 22.5% glucose was started and periodically adjusted to maintain blood glucose concentration at 115–125 mg/dL. Somatostatin (5 μg/kg/min, Bachem) was administered i.v. to inhibit endogenous insulin secretion, and washed donor mouse erythrocytes were infused to maintain blood volume. Blood was collected at the end of the basal and clamp periods to determine EGP (80). A bolus dose of 2-I-[14C] Deoxy-D-glucose (PerkinElmer, 5 μCi) was administered i.v. to measure tissue glucose uptake under steady-state glucose concentrations. Animals were subsequently sacrificed by intraperiarterial administration of pentobarbital. Tissues were rapidly excised and frozen in liquid N2 to estimate tissue glucose uptake (81). Additional arterial blood samples were obtained to monitor hepatic or were stored frozen as plasma for subsequent assays.

Metabolomic analysis. BCAAs, BCKAs, and glutamate were measured using a mass spectrometry-based targeted metabolomics approach. Briefly, 2 overnight extraction protocols were used for the detection of the amino acids (A, acetonitrile-methanol solvent mixture, 50:50% v/v) and the ketoacids (B, methanol-water mixture, 80:20%, v/v), respectively. Aliquots of 25 μL plasma were used for each extraction protocol. Amino acids were also measured in BAT. Tissue was snap frozen in liquid nitrogen and pulverized. A tissue aliquot of about 50 mg was weighted, and the extraction solution A was added to achieve a final concentration of 50 mg/mL. Stable labeled compounds were added to the extraction solutions for internal standard quantitation (Supplemental Table 1). Data were acquired using a Shimadzu Nexera X2 UHPLC system coupled to a Sciex 6500 triple quadrupole mass spectrometer equipped with an electrospray source operating in positive and negative ion mode. BCAAs and glutamate were chromatographically separated using a Waters XBridge BEH Amide column (150 mm × 2.1 mm × 3.5 μm). BCKAs were separated on a Waters XSelect HSS T3 C18 column (150 mm × 2.1 mm × 3.5 μm). The columns were maintained at 40°C. Elution solvents were water with 0.1% formic acid with 10 mM ammonium formate and acetonitrile with 0.1% formic acid. The mobile phases eluted following a linear gradient for both the HILIC and reversed-phased-method, leading to a total of 29 minutes of analysis time per run, including reequilibration (gradient not shown). Data were acquired in multiple-reaction monitoring mode. Details on the MRM conditions used for the analytes of interest are reported in Supplemental Table 1. All positive ions detected were protonated species ([M+H]+). All negative ions were detected as deprotonated species ([M−H]−).

Pharmacokinetic analysis. The PK of LAGIPRA was evaluated in male CD-1 mice following a single i.v. or s.c. dose of 200 nmol/kg. Blood samples were collected over 168 hours, and the resulting individual concentrations from 2 animals/group/time point were used to calculate the reported PK parameters. The mean PK parameters are shown in Figure 4 (Table 2).

Analysis of metabolites and circulating factors. Blood samples were collected on ice before storage of plasma at −80°C. Plasma triglycerides and FFAs were measured using a Hitachi 912 Clinical Chemistry Analyzer (Roche Diagnostics). Insulin (Crystal Chem Inc.), Adiponectin (BioVendor Inc.), IGFBP2 (R&D Systems), and leptin (Crystal Chem Inc.) were measured by ELISA. All samples for a given hormone were assayed in duplicate within a single assay.

Insulin tolerance test. Briefly, on the morning of the procedure, animals were fasted for 4 hours. After which, animals received an intraperitoneal injection of insulin (0.5 U/kg, Humilin R, Eli Lilly and Company). Blood glucose was determined by tail clip at 0, 15, 30, 60, and 120 minutes after injection.

Glucose tolerance test. Briefly, animals were fasted overnight (16 hours). After which, animals received an intraperitoneal injection of glucose (2 g/kg 50% dextrose). Blood glucose was determined by tail clip at 0, 15, 30, 60, and 120 minutes after injection.

Body composition analysis. Body composition of mice was determined using Quantitative Nuclear Magnetic Resonance analysis (ECHO MRI, 3-1 Composition Analyzer; Echo Medical Systems).

RNA isolation and RNA-Seq analysis. Total RNA was isolated from tissues using TRIzol reagent (Invitrogen) and the RNeasy lipid mini kit (Qiagen). RNA quality was confirmed via RNA integrity number (RIN) and using an Agilent RNA ScreenTape Assay with the Agilent 4200 TapeStation (G2991AA). Only samples with RIN values ≥7.9 were chosen for RNA-Seq analysis. Samples were sequenced on an Illumina NovaSeq using paired 150 bp reads. Raw FastQ files were quality trimmed using
cutadapt (cutadapt1.9.1) and aligned using GSnap (v2013-11-27, command line parameters -B 5 -A sam -N 1 -t 8 -s splicessites -quality-proto=sanger -gunzip -sam-multiple-primaries -maxsearch=1000 -npaths=100) to build 38.83 of the mouse genome. Read counts were quantified using a custom Perl script and summarized at the gene level (NCBI m38.p3 annotation). To map reads to genes and obtain gene level expression measures, RNA-Seq data were subjected to a “roll-up” pipeline developed at Eli Lilly and Company. The following rules were applied for the roll-up: (a) exon reads of multiple assays from the same libraries were summed; (b) exons were excluded if more than 80% of samples had less than 10 counts; (c) robust gene level signals across exons of a gene were determined by a robust linear model and were output for each library and each gene; and (d) mean signal of log-transformed gene levels across all samples were median normalized. Read counts were then quantile normalized using a custom R script. Principal component analysis and outlier analysis were applied to the rollup output normalized data to check any potential pattern or outliers among the samples. As no pattern or outlier was detected, the 1-way ANOVA using R software was applied to the normalized data to test the pairwise comparisons among the treatments. For each comparison, the fold changes and P values as well as the FDR (82) were used to adjust for the multiple testing across the genes were reported. Raw and processed data have been deposited within the Gene Expression Omnibus repository (GSE173522).

Statistics. Data are presented as mean ± SEM. Statistical analyses performed included Student’s unpaired 2-tailed t test, 1-way ANOVA or 2-way ANOVA, or Kruskal-Wallis test followed by Holm-Bonferroni or Dunnett’s multiple comparisons test, where appropriate. Differences were considered significant at P<0.05.

Study approval. All experiments were performed in accordance with protocols approved by the Eli Lilly and Co. IACUC.

Author contributions

AKH, JLF, SW, and DLK conducted insulin clamps. MEC conducted insulin clamps and contributed to data interpretation. XR and MD conducted LAGIPRA in vitro characterization and PK analysis. BAD, EPCSF, LD, OC, CCC, DAB, LSOF, RC, and AML performed glucose and insulin tolerance tests and in vivo metabolic analysis, conducted ELISA analyses, and performed RNA isolation. VP and JAW performed metabolomic analysis and contributed data interpretation. KALC and YL were responsible for RNA-Seq analyses. KWS, MPC, REG, JSM, TC, and JAF contributed to data interpretation and writing of the manuscript. WCR was responsible for study conception, data interpretation, and contributed to writing of the manuscript. RJS was responsible for study conception, experimental designs, data analyses and interpretation, and drafting and writing of the manuscript.

Address correspondence to: Ricardo J. Samms or William C. Roell, Lilly Research Laboratories, Eli Lilly and Company, Lilly Corporate Center, Indianapolis, Indiana 46285 USA. Phone: 317.651.7769; Email: Samms_Ricardo_J@lilly.com (RJS). Phone: 317.433.4256; Email: Roell_William_C@lilly.com (WCR).

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