Pharmacological Characterization of Human Incretin Receptor Missense Variants

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ABSTRACT

Glucagon-like peptide 1 (GLP-1) and glucose-dependent insulino tropic polypeptide (GIP) are gut-derived incretin hormones that regulate blood glucose levels. In addition to their widely accepted insulino tropic role, there is evidence that GLP-1 modulates feeding behavior and GIP regulates lipid metabolism, thereby promoting postprandial fat deposition. In this study, we investigated whether naturally occurring polymorphisms in the GLP-1 receptor (GLP-1R) and the GIP receptor (GIP-R) affect the pharmacological properties of these proteins. After transient expression of the receptors in human embryonic kidney 293 cells, basal and ligand-induced cAMP production were assessed by use of luciferase reporter gene assays. Our data reveal that the wild-type GIP-R displays a considerable degree of ligand-independent activity. In comparison, the GIP-R variants C46S, G198C, R316L, and E354Q show a marked decrease in basal signaling that may, at least in part, be explained by reduced cell surface expression. When stimulated with GIP, the C46S and R316L mutants display significantly reduced potency (≥1000 and 25-fold, respectively) compared with wild type. Complementary competition binding assays further demonstrate that the C46S variant fails to bind radioiodinated GIP, whereas all other GIP-R mutants maintain normal ligand affinity. In contrast to the GIP-R, the wild-type GLP-1R lacks constitutive activity. Furthermore, none of the 10 GLP-1R missense mutations showed an alteration in pharmacological properties versus wild type. The extent to which abnormalities in GIP-R function may lead to physiological changes or affect drug sensitivity in selected populations (e.g., obese, diabetic individuals) remains to be further investigated.

The incretin hormones glucose-dependent insulino tropic polypeptide (GIP) and glucagon-like peptide 1 (GLP-1) are homologous peptides released from intestinal enteroendocrine cells in response to food intake. Both hormones are important modulators of metabolic function. In the pancreas, GLP-1 and GIP potentiate nutrient-stimulated insulin secretion and promote the expansion of pancreatic islet mass via induction of β-cell proliferation and survival (Kim et al., 2005; Kim and Egan, 2008). In light of these insulino tropic actions, drugs that mimic or prolong the biological functions of GIP and GLP-1 have attracted considerable attention as treatment options for type 2 diabetes (T2D) (Lovshin and Drucker, 2009). Exendin-4 (Exenatide), a potent long-acting agonist of the GLP-1 receptor (GLP-1R), represents the first incretin-based pharmaceutical to reach the market for the treatment of T2D. Inhibitors of the enzyme dipeptidyl dipeptidase IV, which plays a major role in inactivating both incretin hormones, have also recently been approved as therapeutics for T2D.

Considerable efforts have focused on unraveling additional metabolic functions triggered by the incretins (Kim and Egan, 2008). Accumulating evidence supports that GIP modulates adipocyte metabolism, triggering fat deposition after feeding. Highlighting the physiological relevance of this function, previous studies have shown that targeted disruption of the GIP receptor (GIP-R) in mice results in protection from both diet-induced obesity and insulin resistance (Miyawaki et al., 2002). Consistent with these observations, inhibition of GIP-R signaling using a selective antagonist, or passive immunization against GIP, were both shown to decrease body weight in diet-induced obesity models (Miyawaki et al., 2002). Consistent with these observations, inhibition of GIP-R signaling using a selective antagonist, or passive immunization against GIP, were both shown to decrease body weight in diet-induced obesity models (Miyawaki et al., 2002). Considering the growing interest in the incretin system, the pharmacological characterization of naturally occurring polymorphisms in GIP-R and GLP-1R is of considerable interest.

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ABBREVIATIONS: GIP, glucose-dependent insulino tropic peptide; GLP-1, glucagon-like peptide 1; GLP-1R, glucagon-like peptide 1 receptor; GIP-R, glucose-dependent insulino tropic peptide receptor; GPCR, G protein-coupled receptor; HA, hemagglutinin; CRE, cAMP-responsive element; T2D, type 2 diabetes; ELISA, enzyme-linked immunosorbent assay; HEK, human embryonic kidney; PBS, phosphate-buffered saline.
weight and to protect against glucose intolerance in animals that were fed a high-fat diet (Gault et al., 2007; Fulurija et al., 2008).

GLP-1 also modulates metabolic function, in part, by acting on GLP-1Rs in extrapancreatic tissues (Kim and Egan, 2008). This peptide triggers delayed gastric emptying, which in turn slows the absorption of food, thus delaying the rise in blood glucose levels. In addition, GLP-1 has been shown to inhibit feeding behavior by stimulation of the cognate receptors in the brain. Taken together, the incretin hormones and their receptors contribute at multiple levels to maintaining normal glucose homeostasis and regulating body weight.

Both the GIP-R and the GLP-1R belong to the glucagon subfamily of class B1 G protein-coupled receptors (GPCRs). These seven transmembrane domain proteins, when stimulated with ligand, undergo a conformational change from putative inactive to active conformations, thereby triggering a Gαs-mediated increase in cAMP production (Hoare, 2005). It has been observed with other wild-type and mutant GPCRs that partially active receptor conformations may occur even in the absence of agonist, leading to constitutive, ligand-independent signaling (Kenakin, 2004). Although engineered constitutively active incretin receptors have been generated (Tseng and Lin, 1997; M.B., unpublished data); the extent to which detectable basal signaling is influenced by naturally occurring polymorphic/mutant incretin receptors has not been investigated.

It is well established that missense mutations in GPCRs can result in a variety of pharmacological abnormalities (e.g., alterations in basal and ligand-dependent activity, receptor affinity, expression) which predispose to physiological changes or disease (Seifert and Wenzel-Seifert, 2002). In the current study, we examined the molecular pharmacological consequences of naturally occurring mutations/polymorphisms in the GIP-R and GLP-1R using a series of in vitro assays.

Materials and Methods

Generation of Incretin Receptor Variants. The complementary DNA encoding the GIP-R was obtained from the Missouri S&T cDNA Resource Center (www.cdna.org) and subcloned into pcDNA1.1. The human GLP-1R cDNA was reported previously (Tibaduiza et al., 2001). Single-amino-acid substitutions and a hemagglutinin (HA) tag were introduced into the receptor sequence by use of oligonucleotide-directed site-specific mutagenesis as described previously (Fortin et al., 2009). The nucleotide sequences of all receptor coding regions were confirmed by automated DNA sequencing.

Cell Culture and Transfection. Human embryonic kidney (HEK) 293 cells were grown in Dulbecco’s modified Eagle’s medium (Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine serum, 100 U/ml penicillin G, and 100 μg/ml streptomycin. The cells were maintained at 37°C in a humidified environment containing 5% CO₂.

Luciferase Reporter Gene Assay. Receptor-mediated signaling was assessed by use of a previously described luciferase assay (Fortin et al., 2009). In brief, HEK293 cells were plated at a density of 2000 to 3000 cells/well onto 24-well plates coated with poly-l-lysine, and grown for 18 to 24 h to ~80% confluency. Cells were then transiently transfected by use of Lipofectamine reagent (Invitrogen) with cDNAs encoding 1) a GPCR (or empty expression vector), 2) a cAMP responsive element-luciferase reporter gene (CRElox-luc), and 3) β-galactosidase to enable correction for interwell variability in transfection efficiency and cell survival. Forty-eight hours after transfection, cells were incubated for 6 h with or without the appropriate peptide ligand (American Peptide Company Inc., Sunnyvale, CA) in serum-free medium. After agonist treatment, the medium was gently aspirated, the cells were lysed, and luciferase activity was measured by use of Steadylute reagent (PerkinElmer Life and Analytical Sciences, Waltham, MA). A β-galactosidase assay was then performed after adding the enzyme substrate, 2-nitrophenyl β-d-galactopyranoside. After incubation at 37°C for 30 to 60 min, substrate cleavage was quantified by measurement of optical density at 420 nm using a SpectraMax microplate reader (Molecular Devices, Sunnyvale, CA). Corresponding values were used to normalize the luciferase data.

Assessment of Receptor Expression with Use of ELISA. The surface expression levels of the HA-tagged GIP-Rs were assessed by use of a previously described approach (Shinyama et al., 2003). In agreement with earlier reports (Lee et al., 1994; Qi et al., 1997), two independent predictor tools (http://bmbpcu36.leeds.ac.uk/prot_analysis/Signal.html and http://www.cbs.dtu.dk/services/) supported the presence of a signal sequence in the GIP-R extracellular domain that is cleaved during receptor maturation. An HA tag was thus inserted immediately downstream of the putative 24-amino acid GIP-R signal peptide (shown in Fig. 1B). HEK293 cells grown in 96-well clear Primaria plates (BD Biosciences Discovery Labware, Bedford, MA) were transiently transfected with increasing amounts of either pcDNA1.1 or a cDNA encoding the HA-tagged GIP-R. Forty-eight hours after transfection, the cells were washed once with phosphate-buffered saline (PBS), pH 7.4, and fixed with 4% paraformaldehyde in PBS for 10 min at room temperature. After washing with 100 mM glycine in PBS, the cells were incubated for 30 min in blocking solution (PBS containing 20% bovine serum). A horseradish peroxidase-conjugated antibody directed against the HA epitope tag (Roche; clone 3F10, monoclonal, 1:500 in blocking buffer) was then added to the cells. After 1 h, the cells were washed five times with PBS. Fifty microliters per well of a solution containing the peroxidase substrate BM-blue (3′,3′,5′,5′-tetramethylbenzidine; Roche Applied Science, Indianapolis, IN) was then added. After incubation for 30 min at room temperature, conversion of this substrate by antibody-linked horseradish peroxidase was terminated by adding 2.0 M sulfuric acid (50 μl/well). Results were quantified by measuring light absorbance at 450 nm.

Radioligand Binding Studies. HEK293 cells were plated at a density of 30,000 cells/well onto 24-well plates coated with poly-l-lysine, and grown for 18 to 24 h to ~80% confluency. Cells were then transiently transfected by use of lipofectamine with receptor cDNA (100 ng/well) and grown for an additional 18 to 24 h. Whole cell binding studies were initiated by washing cells twice with cold (4°C) assay buffer (Dulbecco’s modified Eagle’s medium with 0.1% bovine serum albumin and 15 mM HEPES), followed by addition of the same media with 125I-GIP (PerkinElmer Life and Analytical Sciences) and varying concentrations of unlabeled GIP. After an 8-h incubation period at 4°C, the cells were washed twice with cold assay buffer and solubilized in 0.1 N NaOH. The lysates were then counted by use of a Packard Cobra Quantum γ-counter to determine cell-associated radioactivity.

Data and Statistical Analysis. GraphPad Prism software version 5.0 (GraphPad Software Inc., San Diego, CA) was used for sigmoidal curve fitting. Half-maximal effective concentrations (EC₅₀ values) were calculated as an index of ligand potency, and half-maximal inhibitory concentrations (IC₅₀ values) were calculated as an index of receptor binding affinity. pEC₅₀ basal activity and surface expression values for each of the mutants were compared with the corresponding control value at the wild-type receptor by use of one-way analysis of variance followed by Dunnett’s post test (GraphPad INSTAT software).
Results

Human Incretin Receptor Variants. Receptor constructs containing naturally occurring missense mutations of the human GLP-1R (P7L, R20K, R44H, R131Q, G168S, F260L, A316G, A316T, S333C, and R421C) and human GIP-R (C46S, R136W, G198C, A207V, L262V, R316L, F260L, A316G, A316T, S333C, and R421C) were generated for investigation. The position of each amino acid substitution is illustrated in cartoons of the GLP-1R and GIP-R (Fig. 1, A and B, respectively). Each incretin receptor variant appeared in the NaVa (Natural Variants) database that catalogs known human GPCR polymorphisms (frequency >1%), as well as rarer mutants (Kazius et al., 2008). As outlined in the discussion, three GIP-R variants have been described previously in the literature (Kubota et al., 1996; Almind et al., 1998). Site-directed mutagenesis was used to introduce amino acid substitutions corresponding to the receptor variants. Each of the mutant receptor constructs or corresponding wild-type proteins were expressed in HEK293 cells and pharmacologically characterized.

Missense Variants of the GLP-1R Exhibit Normal Basal and Agonist-Induced Signaling. Basal signaling in cells expressing the wild-type GLP-1R (assessed by use of a cAMP-responsive luciferase construct) was indistinguishable from that observed in cells transfected with the empty expression vector, pcDNA1.1 (data not shown). This observation confirms that the GLP-1R lacks constitutive activity. In addition, none of the 10 GLP-1R variants showed a significant level of basal signaling (Table 1).

Agonist-induced GLP-1R function was assessed by use of two structurally related agonists, GLP-1 and exendin-4 (illustrated for the wild-type receptor and representative variants in Fig. 2). At each mutant receptor, both peptides demonstrated potency and efficacy values that are comparable with wild type (Table 1).

Selected GIP-R Variants Show Altered Basal and/or GIP-Mediated Activity. Basal and GIP-induced signaling was examined at each GIP-R isoform (Fig. 3). In contrast to the GLP-1R, the wild-type GIP-R showed constitutive activity (~25.0 ± 4.8% of the GIP-induced maximum) that markedly exceeded control values (determined by use of vector-transfected cells). Four GIP-R variants, C46S, G198C, R316L, and E354Q, showed a significant reduction in basal activity (Table 2). Of these functionally abnormal receptors, two also showed a marked decrease in GIP potency. The C46S variant showed a greater than 1000-fold reduction, whereas R316L had a ~25-fold decrease in GIP potency compared with that at the wild-type GIP-R. In contrast, the EC50 values for GIP at the R136W, G198C, A207V, L262V, E354Q, and E463Q mutants were comparable with the reference value at the wild-type receptor (Table 2).

Impaired Binding Affinity of GIP at the C46S GIP-R Variant. To complement the functional studies of GIP-R mutants, we evaluated the affinity of GIP at each receptor variant by radioligand competition binding assays (Fig. 4). At each mutant receptor, both peptides demonstrated potency and efficacy values that are comparable with wild type (Table 2). At each receptor variant, both peptides demonstrated potency and efficacy values that are comparable with wild type (Table 2).

### Table 1

<table>
<thead>
<tr>
<th>Receptor</th>
<th>GLP-1</th>
<th>Exendin-4</th>
<th>Basal Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EC50</td>
<td>pEC50a</td>
<td>EC50</td>
</tr>
<tr>
<td>GLP-1 Variant</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>hGLP-1R</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P7L</td>
<td>1.8</td>
<td>11.75 ± 0.04</td>
<td>1.3</td>
</tr>
<tr>
<td>R20K</td>
<td>1.7</td>
<td>11.73 ± 0.07</td>
<td>1.1</td>
</tr>
<tr>
<td>R44H</td>
<td>2.0</td>
<td>11.70 ± 0.07</td>
<td>1.6</td>
</tr>
<tr>
<td>R131Q</td>
<td>1.8</td>
<td>11.74 ± 0.09</td>
<td>1.1</td>
</tr>
<tr>
<td>R316L</td>
<td>1.6</td>
<td>11.79 ± 0.09</td>
<td>1.3</td>
</tr>
<tr>
<td>F260L</td>
<td>2.2</td>
<td>11.65 ± 0.11</td>
<td>1.4</td>
</tr>
<tr>
<td>A316G</td>
<td>2.0</td>
<td>11.58 ± 0.07</td>
<td>2.4</td>
</tr>
<tr>
<td>A316T</td>
<td>2.6</td>
<td>11.70 ± 0.11</td>
<td>1.6</td>
</tr>
<tr>
<td>S333C</td>
<td>2.2</td>
<td>11.66 ± 0.10</td>
<td>1.7</td>
</tr>
<tr>
<td>R421C</td>
<td>2.6</td>
<td>11.58 ± 0.11</td>
<td>1.7</td>
</tr>
</tbody>
</table>

a No significant difference vs. wild type.

b Percentage of the corresponding GLP-1-induced maximum.
respectively. Data represent the mean in independent experiments, each performed in triplicate. Representative activity were 3.51 values for basal and GLP-1/exendin-4-induced maximum luciferase ac-

induced maximal stimulation (A or B) at the wild-type GLP-1R. Average activity values were normalized relative to the GLP-1- or exendin-4-

activity was quantified as described under Materials and Methods. All activity values were normalized relative to the GIP-stimulated maximum at the wild-type GIP-R. Average values for basal and GIP-induced maximum luciferase activity were 1.50 ± 0.22 × 10^6 and 5.62 ± 0.70 × 10^6 cps, respectively. Data represent the mean ± S.E.M. from at least three independent experiments, each performed in quadruplicate.

Fig. 2. All GLP-1R variants show a pharmacological response to GLP-1 and exendin-4 that is similar to wild type. HEK293 cells were transiently transfected with a receptor-encoding cDNA and a CRE-Luc reporter gene construct. Forty-eight hours after transfection, cells were stimulated for 4 h with media containing either no peptide (basal) or increasing concentrations of GLP-1 (A) or exendin-4 (B). After stimulation, luciferase activity was quantified as described under Materials and Methods. All activity values were normalized relative to the GLP-1- or exendin-4-induced maximal stimulation (A or B) at the wild-type GLP-1R. Average values for basal and GLP-1/exendin-4-induced maximum luciferase activity were 3.51 ± 0.62 × 10^6 and 2.38 ± 0.34 × 10^6/2.34 ± 0.21 × 10^6 cps, respectively. Data represent the mean ± S.E.M. from at least three independent experiments, each performed in triplicate. Representative variants are shown.

GIP-R Variants with Decreased Basal Activity Also Show Reduced Cell Surface Expression. Reduced basal activity of several GIP-R variants (C46S, G198C, R316L, and E354Q; Fig. 3) was confirmed in experiments where ligand-independent signaling was measured after transfecting cells with increasing concentrations of respective receptor cDNAs (Fig. 5A). In a parallel experiment using the same transfection protocol, receptor expression levels at the cell surface were determined by ELISA (Fig. 5B). These studies revealed that, at each cDNA level, receptors with reduced basal signal-

Fig. 3. Selected GIP-R mutations alter GIP-induced signaling. HEK293 cells were transiently transfected with the empty vector pcDNA1.1 or a receptor-encoding cDNA, together with a CRE-Luc reporter gene con-

struct. Forty-eight hours after transfection, cells were stimulated for 6 h with media containing either no peptide (basal) or increasing concentrations of GIP. After stimulation, luciferase activity was quantified as described in Materials and Methods. All activity values were normalized relative to the GIP-stimulated maximum at the wild-type GIP-R. Average values for basal and GIP-induced maximum luciferase activity were 1.50 ± 0.22 × 10^6 and 5.62 ± 0.70 × 10^6 cps, respectively. Data represent the mean ± S.E.M. from at least three independent experiments, each performed in quadruplicate.

Discussion

The related peptides, GIP and GLP-1, play important physiological roles in maintaining blood glucose homeostasis, most notably by potentiating glucose-stimulated insulin secretion by pancreatic β-cells (Kim and Egan, 2008; Lovshin and Drucker, 2009). These peptides have additional periph-
TABLE 2
Pharmacological properties of wild-type vs. mutant GIP-Rs
All values represent the mean ± S.E.M. from at least three independent experiments. Functionally abnormal variants are highlighted.

<table>
<thead>
<tr>
<th>Receptor</th>
<th>EC50 (pM)</th>
<th>pEC50</th>
<th>GIP binding Kd (nM)</th>
<th>Basal Activityb</th>
<th>Surface Expressionc</th>
<th>Slopec</th>
</tr>
</thead>
<tbody>
<tr>
<td>hGIP-R</td>
<td>0.9</td>
<td>12.06 ± 0.11</td>
<td>3.1 ± 0.6</td>
<td>100</td>
<td>100</td>
<td>1.0</td>
</tr>
<tr>
<td>C46S</td>
<td>&gt;1000</td>
<td>&lt;7</td>
<td>&gt;1000</td>
<td>34 ± 4</td>
<td>31 ± 6</td>
<td>1.1</td>
</tr>
<tr>
<td>R136W</td>
<td>1.8</td>
<td>11.70 ± 0.19</td>
<td>3.9 ± 0.6</td>
<td>102 ± 1</td>
<td>102 ± 8</td>
<td>0.9</td>
</tr>
<tr>
<td>G198C</td>
<td>2.0</td>
<td>11.75 ± 0.12</td>
<td>1.7 ± 0.8</td>
<td>22 ± 4</td>
<td>56 ± 9</td>
<td>0.3</td>
</tr>
<tr>
<td>A207V</td>
<td>1.0</td>
<td>11.97 ± 0.05</td>
<td>2.6 ± 0.9</td>
<td>104 ± 10</td>
<td>99 ± 5</td>
<td>1.1</td>
</tr>
<tr>
<td>L262V</td>
<td>1.0</td>
<td>12.06 ± 0.12</td>
<td>2.9 ± 0.4</td>
<td>99 ± 11</td>
<td>97 ± 13</td>
<td>1.0</td>
</tr>
<tr>
<td>R316L</td>
<td>24.3</td>
<td>10.75 ± 0.20</td>
<td>2.7 ± 0.4</td>
<td>3 ± 1</td>
<td>37 ± 7</td>
<td>0.01</td>
</tr>
<tr>
<td>E254Q</td>
<td>0.7</td>
<td>12.22 ± 0.11</td>
<td>2.2 ± 0.5</td>
<td>15 ± 6</td>
<td>60 ± 10</td>
<td>0.2</td>
</tr>
<tr>
<td>E463Q</td>
<td>1.3</td>
<td>11.96 ± 0.13</td>
<td>2.7 ± 0.9</td>
<td>100 ± 11</td>
<td>99 ± 5</td>
<td>1.0</td>
</tr>
</tbody>
</table>

a Value significantly different (P < 0.01) vs. wild-type GIP-R value.
b Percentage of basal signaling activity of the wild-type GIP-R.
c Percentage of wild type GIP-R surface expression.
d Slope of regression line in Fig. 5C.

Fig. 4. Effect of GIP-R mutations on GIP binding affinity. [125I-GIP radioligand binding with increasing concentrations of unlabeled GIP was evaluated in HEK293 cells transiently expressing either the wild-type or a mutant GIP-R. The cells were incubated in the presence of radioligand with wild-type values. Mutation-induced abnormalities in agonist potency may be triggered by two distinct mechanisms: 1) alteration of the hormone binding site and/or 2) defective GPCR transitioning from the inactive to the active receptor state (the conformation triggering G protein activation) (Beinborn et al., 2004).

The C46 substitution is found in the N terminus of the GIP-R, a domain that plays an important role in ligand binding (Parthier et al., 2007). It is thus likely that the observed decrease in potency and the absence of radioligand binding to the C46S variant (Table 2) is due to a mutation-induced alteration of the hormone binding domain. This conclusion is supported by analysis of the recently obtained crystal structure of the GIP-R extracellular domain bound to GIP. Experimental evidence from this study suggests that three conserved disulfide bridges, including a link between C46 and C70, stabilize the secondary structure of the extracellular domain (Parthier et al., 2007). Furthermore, mutation of homologous cysteine residues in other class B1 GPCRs have been shown to disrupt ligand affinity (Lee et al., 1994; Gaudin et al., 1995; Qi et al., 1997; Lisenbee et al., 2005).

In contrast to C46S, the other GIP-R polymorphism that decreases agonist potency (R316L) is found in the third intracellular loop. Because this receptor region is far removed from the ligand binding domains, it is unlikely that the reduced GIP potency observed at the R316L variant results from a direct change in the hormone docking site. Consistent with this conclusion, the R316L mutant maintains normal affinity for radiiodinated GIP (which reflects the initial step of ligand-receptor interaction) despite reduced agonist potency (a measure of subsequent ligand-induced receptor activation). It is noteworthy that previous structure-function studies on the related GLP-1R and parathyroid hormone receptor revealed that important G protein coupling determinants localize in the N-terminal section of the third intracellular loop of these receptors (i.e., the region where R316L is found in the GIP-R) (Huang et al., 1996; Mathi et al., 1997). It is thus probable that the reduction in GIP potency at the R316L isoform reflects an altered ability of this variant to couple and/or activate stimulatory G proteins.

Our studies revealed normal GIP potency and affinity for the G198C mutant (Table 2). It is noteworthy that this result contrasts with an earlier study that reported lower potency for this GIP-R variant relative to wild type (Kubota et al., 1996). The basis for this discrepancy is not clear, but it is possible that the divergent findings are at least in part ex-
receptor-mediated function may only be detectable when
of possible mutation-induced changes. Some alterations in
characterization (including the choice of cells for cDNA
explained by differences in methodologies used for receptor
crystalline of our current findings, this prior study revealed that
of the dopamine D1R and D2R (Al-Fulaij et al., 2008). Remi-
functional abnormalities of GIP-R variants, including the
pharmacologically distinct from wild type) contrasts with our
in vitro findings provide valuable insight into the potential of
mutations to affect receptor function (Seifert and Wenzel-
expressed in triplicate.
illustrated in methods used for receptor
that the wild-type or a mutant
receptor property.
Four GIP-R variants (C46S, G198C, R316L, and E354Q) are
characterized by a significant reduction in ligand-indepen-
dependent signaling relative to wild type (Fig. 5, Table 2). For
one of these mutants (C46S), this functional change appears
to be largely accounted for by reduced cell surface expres-
sion, whereas additional factors may underlie the de-
creased basal activity of the G198C, R316L, and E354Q
variants. As a contributing mechanism, mutation-induced
structural changes may shift the putative equilibrium be-
tween active and inactive receptor conformations (Lefkow-
itz et al., 1993) and/or may alter G protein-receptor inter-
action (as discussed above for the R316L mutant). Given
that the G198C and E354Q substitutions are localized
outside the intracellular receptor portion (in EC loop I and
transmembrane domain VI), it is possible that these mu-
tations induce structural changes that primarily shift the
receptor equilibrium and thereby indirectly compromise G
protein interaction.
Loss of function in the GIP-R could provide a potential
mechanism for altered glucose homeostasis or fat deposition
(Miyawaki et al., 2002; Gault et al., 2007; Fulurija et al.,
mechanism for altered glucose homeostasis or fat deposition
protein interaction.
that the G198C and E354Q substitutions are localized
outside the intracellular receptor portion (in EC loop I and
transmembrane domain VI), it is possible that these mu-
tations induce structural changes that primarily shift the
receptor equilibrium and thereby indirectly compromise G
protein interaction.
the E354Q polymorphism (a variant which in our hands
showed reduced basal activity) had a decreased serum C-
peptide concentration (an index of insulin secretion) (Almind
study reported that glucose-tolerant subjects homozygous for
the E354Q polymorphism (a variant which in our hands
showed reduced basal activity) had a decreased serum C-
peptide concentration (an index of insulin secretion) (Almind
et al., 1998). This abnormality was observed under fasting
conditions and after an oral glucose load, relative to subjects
with the wild-type GIP-R. Future efforts will explore whether
functional abnormalities of GIP-R variants, including the
E354Q polymorphism, contribute to metabolic phenotypes.
The relatively high rate of mutation-induced functional
changes in the GIP-R (four of the eight known variants were
pharmacologically distinct from wild type) contrasts with our
parallel analysis of the GLP-1R where none of the 10 vari-
ants that were investigated showed detectable abnormalities.
On this background, it is noteworthy that the GIP-R is
constitutively active (Figs. 3 and 5). There is only one prior
report in the literature suggesting that the GIP-R has a low
degree of constitutive activity (Almind et al., 1998). Our
ability in the current study to readily detect a pronounced
elevation in GIP-R basal activity is likely explained by the
sensitivity of the luciferase-based system that was used to
assess receptor-mediated signaling. For the broader group of
class B1 GPCRs (i.e., the secretin-glucagon family), there are few reports of significant ligand-independent
signaling of unmodified wild-type receptors (Seifert and
Wenzel-Seifert, 2002; Hoare et al., 2008). Our demonstration
of GIP-R constitutive activity provided the basis on
which to define the effects of specific missense mutations
on this receptor property.
illustrating this limitation, the current study is the first to
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Allosteric ligands for the corticotropin releasing factor type 1 receptor modulate conformational states involved in receptor activation. Mol Pharmacol 73:1371–1376.


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