Three intragenic suppressors of a GTPase-deficient allele of GNAS associated with McCune–Albright syndrome

Kyle Turcic, Raquel Tobar-Rubin, Daniela Janevska, Julie Carroll, Eraj Din, Rebecca Alvarez, Jennifer Haick and Robin Pals-Rylaarsdam

Department of Biological Science, Benedictine University, 5700 College Road, Lisle, Illinois 60532, USA

Abstract

Gain-of-function mutations in heterotrimeric G-protein α subunits are associated with a variety of human diseases. McCune–Albright syndrome (MAS) is caused by mutations in GNAS, the gene encoding Gs. Alterations at Arg201 significantly reduce the GTPase activity of the protein, rendering it constitutively active. In this study, we have constructed a library of random mutations in a constitutively active yeast GPA1 gene carrying a mutation homologous to the McCune–Albright allele (Arg297His). Intragenic suppressors found at sites with homology to the human Gs protein were tested for their ability to suppress the constitutive activity of an Arg201His mutation in Gs. Three intragenic suppressors, at Phe142, Arg231, and Leu266, were able to suppress elevated basal cAMP responses caused by Arg201His when expressed in HEK293 cells. A range of amino acid substitutions was introduced at each of these sites to investigate the chemical requirements for intragenic suppression. The ability of Gs proteins carrying the suppressor mutations alone to mediate receptor-induced cAMP production was measured. These results offer potential sites on Gs that could serve as drug targets for MAS therapies.

Key Words

- McCune–Albright syndrome
- G-protein
- cAMP
- suppressor mutation

Introduction

McCune–Albright syndrome (MAS) was first described in the 1930s as a trio of associated symptoms which included café-au-lait skin hyperpigmentation, precocious puberty, and polyostotic fibrous dysplasia of bone (McCune 1936, Albright et al. 1937). A wide variety of hyperfunctioning endocrine disorders may accompany the classic triad of symptoms, including growth hormone excess, renal phosphate wasting, and Cushing’s syndrome (Dumitrescu & Collins 2008). This widely varied disease is caused by mosaicism for mutations in GNAS, the gene encoding the Gsα subunit of heterotrimeric G-proteins (Gs; Weinstein et al. 1991). Substitution of arginine 201 with another amino acid reduces GTP hydrolysis, causing constitutive activity of the protein and its signaling pathways. Histidine and cysteine substitutions are most common (Weinstein et al. 1991, Schwindinger et al. 1992, Lumbroso et al. 2004), although serine (Candeliere et al. 1997) and glycine (Riminucci et al. 1999) have also been found at codon 201 in MAS isolates.

The endocrine disorders associated with MAS are managed with drugs. For example, the full onset of puberty in MAS patients is delayed until the patient...
approaches a normal age for puberty by using aromatase inhibitors (Feuillan et al. 2007) and/or tamoxifen (Eugster et al. 2003). Fibrous dysplasia often requires repeated surgeries for repair of broken or deformed long bones, or for correction of facial deformities (Collins 2006, Stanton et al. 2012). Clinical trials using bisphosphonates showed a reduction in bone pain (Liens et al. 1994), but no effectiveness in preventing the spread or development of fibrous lesions in children with MAS (Plotkin et al. 2003, Chan & Zacharin 2006). Clearly, more effective therapies for the bone dysplasias would benefit MAS patients.

One experimental approach to identify targets for rational drug design is to identify intragenic suppressors of mutations associated with a disease, and to analyze the suppressors to gain structural information about the potential drug target. The yeast *Saccharomyces cerevisiae* is a powerful genetic system for approaching the identification of suppressor mutations, and in particular suppressors of activating mutations in Gs subunits (Apanovitch et al. 1998, Ooms et al. 2006). A great deal is known about the structure of Gs, which aids in the interpretation of the suppressor mutations and their potential as drug targets (Sunahara et al. 1997, Tesmer et al. 1997, Rasmussen et al. 2011). Our laboratory previously undertook a pilot study which randomly mutated a constitutively active allele of GPA1 to search for intragenic suppressors of R297H, homologous to the MAS mutation in Gs. The study identified one site for intragenic suppression, located in the GTP-binding site of the G-protein (Ooms et al. 2006). This report describes a full-scale screen for more intragenic suppressors of constitutively active human Gs, and demonstrates three unique locations on the protein that can reverse the constitutive activity: a residue in the helical domain, one within Switch II, and one that interacts with residues of Switch III.

**Materials and methods**

**Yeast strains and plasmids**

RM20 (Medici et al. 1997) was a gift from Gianfranco DiSegni. It is a *gpa1*Δ *ste2*Δ haploid strain (MATa *gpa1*Δ *ste2*Δ fis1::FUS1-lacZ[TRP1] *ura3*::Scr1 *ade2-1* his3-11,5 *leu2-3,112*) containing the plasmid pYS212 (2u-*URA3*-TPI promoter; R&D Systems, Minneapolis, MN, USA) carrying the *GPA1* gene. The *gpa1*Δ deletion in this strain is deleted for the 0.84 kb HindIII–SphI fragment of the coding region. The WT yeast strain MHYS00 (MATa his3Δ200, *leu2-3,112* *ura3*- *S2* *lys2*::801 *tpi1*-1; Chen et al. 1993) came from Mark Hochstrasser’s Laboratory, who also supplied plasmid p413GPD (CEN, HIS3, GPD promoters; Mumberg et al. 1995), used to express an EcoRI fragment encoding GPA1 or its mutated allele R297H in yeast cells. The GPA1 alleles were engineered to contain restriction sites for Ncol at nucleotide 151 and Hpal at nucleotide 1620 (Ooms et al. 2006).

**Yeast culture**

YPD-rich yeast medium contains 1% yeast extract, 2% BactoPeptone, and 1% glucose. SD dropout medium contains 0.67% yeast nitrogen base without amino acids, 1% glucose, supplemented with the appropriate amino acids, adenine, and uracil. Fluoro-orotic acid (FOA) plates are SD-His dropout plates supplemented with 0.2 mg/ml uracil and 1% 5-FOA. All yeast incubations were carried out at 30°C.

**Library construction**

ACGT, Inc. (Wheeling, IL, USA) constructed a library of random mutations in the *S. cerevisiae* GPA1 gene using degenerate PCR. Plasmid p413GPD-GPA1 (R297H) was used as the template for a PCR using the Diversity PCR Random Mutagenesis Kit (Clontech) with 8 mM MnSO₄, 2 mM dGTP, and 10 mM of each primer (forward, 5′-CATACATAACGAGGGTGTATAAACATGA-3′ and reverse, 5′-CTAAACACTAACTGAATTACAGTCCCTTC-3′) and 10 ng/µl template DNA. Cycling conditions were 94°C for 30 s, then 25 cycles of 94°C for 30 s, and 68°C for 90 s. The cycle ended with 68°C for 1 min and a 4°C soak. The amplified PCR product was separated on a 0.8% agarose gel and showed a single band at the expected 1.4 kb. The product was then ligated into the pGEM-T Easy Vector (Promega) and transformed into Top10 chemically competent cells (ACGT, Inc.). The transformed cells were spread onto Luria broth (LB) plates supplemented with ampicillin and X-gal and grown overnight. Plasmid DNA was isolated from 39 clones using the Qiaprep Spin Miniprep Kit (Qiagen). The insert DNA was sequenced from both strands using the following primers in addition to the M13-F and M13-R universal primers: GPA1_F1, 5′-ACCTTGAATTTGGACGAGC-3′; GPA1_F2, 5′-TCGAAGGAATTACAGTCCCTTC-3′; GPA1_R1, 5′-ACCTTGAATTTGGACGAGC-3′; and GPA1_R2, 5′-CATACATAACGAGGGTGTATAAACATGA-3′. Sequence data from both strands was aligned and a consensus sequence generated. The sequence of each mutant clone was compared with the reference sequence to determine the number of amino acid changes per clone. Of the 39 clones, 12 exhibited no changes, 22 clones exhibited one to three amino acid changes, and five
exhibited four to seven amino acid changes. Thirteen large (150 mm) plates with ~600 clones/plate were delivered from ACGT, Inc. These colonies were pooled, and the plasmid DNA was purified directly from the colonies without outgrowth using an E.Z.N.A. Plasmid Maxi Kit from Omega Biotek (Norcross, GA, USA). The mutated \( \text{GPA1} \) was excised from \( \text{pGEM-T} \) using NcoI and HpaI, subcloned into pRS413GPD-GPA1, and transformed into NEB 5-alpha \( \text{Escherichia coli} \). Approximately, 55 000 colonies were pooled, grown in 500 ml LB/ampicillin for 6 h at 37 °C with shaking, and then either frozen back as library stocks or used in a plasmid maxi-prep for yeast transformation. Since this expression vector library was derived from 7800 unique clones, we estimate that each clone is represented seven times in our library, and that the expression library, like the \( \text{pGEM} \) collection, has only 70% of its clones with amino acid substitutions.

**Library screen**

Yeast were transformed using the Frozen-EZ Yeast Transformation II Kit (Zymo Research, Irvine, CA, USA) and grown on SD-His plates to select for uptake of the library plasmid. Colonies were replica plated to FOA plates, which selected for loss of the URA3-marked plasmid carrying the WT GPA1 allele. Growth of yeast on FOA was followed for 4–6 days. FOA-resistant clones were grown overnight in SD-His liquid medium, the plasmids isolated using the Zymoprep Yeast Plasmid Miniprep II Kit (Zymo Research, Irvine, CA, USA), and then transformed into NEB 5-alpha high efficiency competent \( \text{E. coli} \) (New England Biolabs, Ipswich, MA, USA). Plasmids were purified from \( \text{E. coli} \) using the Wizard Plus SV Minipreps DNA Purification System (Promega), re-transformed into RM20 yeast plated on SD-His medium. Colonies from these transformations were patched onto FOA plates to retest for plasmid-dependent FOA resistance. Plasmids that supported RM20 growth on FOA at this point were sequenced (ACGT, Inc.) by the same primers used in the mutagenic PCRs.

**Site-directed mutagenesis**

Site-directed mutagenesis of pCDNA3.1+GNAS\( _L \) was carried out using the Agilent Quick-Change XL Mutagenesis Kit (Agilent, Santa Clara, CA, USA). Sense primer sequences for all mutagenesis reactions are shown in Supplementary Table 1, see section on supplementary data given at the end of this article. Antisense primers were the exact complement of the sequences above. Primers were synthesized by Life Technologies. Each mutagenic primer also introduced a new restriction endonuclease site into the sequence while changing the amino acid for only the mutated site. Miniprep DNA was digested with the appropriate restriction enzyme(s) and analyzed by agarose gel electrophoresis. All mutagenesis was confirmed by dideoxy sequencing in both directions performed by ACGT, Inc.

**Mammalian cell culture**

HEK293 cells (ATCC, Manassas, VA, USA) passages 5–17 were maintained in DMEM supplemented with 10% v/v fetal bovine serum, 1% v/v penicillin–streptomycin, and 1.25 mg/ml amphotericin B at 37 °C in a humidified 5% CO\( _2 \) incubator. All cell culture reagents were purchased from Life Technologies.

**Transfection**

Typically, 60 mm plates at 50–60% confluence were transfected with plasmid DNA using JetPRIME reagent (Polyplus-Transfection, Inc., New York, NY, USA). Up to 5 ml of media covered each plate, and cells were transfected in a total volume of 400 \( \mu \)l containing plasmid, 10 \( \mu \)l JetPRIME reagent, and the remaining volume JetPRIME buffer. The transfection mixture was dripped directly onto the culture medium, and no refeeding occurred. 24 h post-transfection, the cells were replated for parallel western blot and signaling assays. Assays for protein expression and second-messenger production were carried out 48 h post-transfection.

**cAMP assays**

The cells were replated 24 h post-transfection by trypsinization followed by resuspension of the cells in 7 ml of culture medium. 100 \( \mu \)l of cells were plated in each of 16 wells of a 96-well culture plate. The remaining cells were allowed to reattach to the 60 mm culture dish. 48 h post-transfection, the culture medium was removed, and the cells were exposed to 100 \( \mu \)l of varying doses of human chorionic gonadotropin (hCG) or forskolin (1 \( \mu \)M) in serum-free DMEM supplemented with 1 mM 3-isobutyl-1-methylxanthine. Following 15 min of drug treatment at 37 °C, cells were lysed by adding 100 \( \mu \)l 0.2 M HCl. Lysates were frozen at −20 °C until protein expression was verified by western blot. Levels of cAMP in the lysates were assayed using a Direct cAMP ELISA Kit (Enzo Life Sciences, Plymouth Meeting, PA, USA) and reported as a percentage of the maximal forskolin-stimulated cAMP levels to control for differences in transfection efficiency from experiment to experiment.
Western blot

Whole-cell lysates of transfected cells were prepared by removing the transfected cells from the culture plate in PBS, pelleting the cells with a low-speed centrifuge, and lysing the pellet directly with 100 μl of Lammeli sample buffer in boiling water for 2 min. Proteins were sonicated at 50% power in a Misonix XL-2000 sonicator for 2–15 s exposures, spun in at 16000 g in a microcentrifuge, and then analyzed by western blot immediately or stored in a −20 °C freezer until analysis. 20 μl of cell lysate was applied to each well of 4–20% Tris-tricine gradient gel (Expedeon, San Diego, CA, USA), and proteins were separated by electrophoresis and transferred to nitrocellulose. Following 30 min of blocking with LI-COR blocking buffer (LI-COR Biosciences, Lincoln, NE, USA), mouse anti-human Gαs MAB12 (Santa Cruz Biotechnology) was diluted 1:2000 in LI-COR blocking buffer and incubated with the membrane for 1–2 h at room temperature or overnight at 4 °C with rocking. The membrane was washed five times for 5 min each with PBS, and then incubated 1–2 h at room temperature or overnight at 4 °C with rocking with a 1:10,000 dilution of IRDye 800CW Goat anti-Mouse IgG (LI-COR Biosciences). The membranes were washed five times for 5 min each in PBS before detection. The blots were imaged with a LI-COR Odyssey Fc for infrared detection of the reactive bands.

Statistical analysis and curve fitting

GraphPad Prism (GraphPad Software, LaJolla, CA, USA) was used for curve fitting and calculation of EC50 values. Assays for basal and receptor-stimulated cAMP were analyzed by ANOVA followed by Fisher’s least significant difference analysis when the ANOVA indicated evidence of a difference among the means.

Results

Our goal in these experiments was to identify secondary mutations in Gs that can compensate for, or suppress, the constitutive activity of the R201H mutation associated with MAS. We designed a yeast system where the yeast α subunit Gpa1 carried the homologous mutation, R297H, along with a library of other random mutations in the gene. Yeast cell division is arrested upon release of bg from the Gpa1 α subunit; thus, cells carrying only a constitutively active Gpa1-like R297H undergo cell-cycle arrest. The cells were maintained in a dividing state by providing a second WT copy of Gpa1 on a URA3-marked plasmid. Transfer to 5-FOA plates selected for loss of the WT plasmid, as Ura3 metabolizes 5-FOA to a toxin. The remaining GPA1 allele supported colony formation only if the random mutations reversed the constitutive activity of R297H, allowing binding to βγ and cell cycle progression (Ooms et al. 2006). Plasmids were isolated from 166 colonies and the GPA1 coding region was sequenced in its entirety. Mutations that reverted histidine back to arginine and that were not homologous to the human Gs were not analyzed further.

Twelve of the unique mutations that suppressed the constitutive activity of GPA1-R297H were located at sites
conserved with the human Gs protein. Site directed mutagenesis was used to introduce these 12 mutations into the human GNAS cDNA which also carried the MAS R201H mutation. Eleven of these mutant isoforms were successfully expressed in HEK293 cells at levels similar to WT and the R201H mutant (Fig. 1a); substitution of H220 with an asparagine residue consistently produced no protein detectable by western blot. Levels of basal cAMP in transfected cells were measured and reported as a percent of the forskolin response to control for changes in cell number per well from experiment to experiment. In agreement with previous studies, transfection of WT Gs resulted in a low level of basal cAMP while expression of Gs-R201H strongly increased the basal cAMP levels in the cells. While most of the mutations that suppressed the constitutively active GPA1-R297H had no effect on suppressing the human MAS mutation, three sites were identified that exhibited basal levels of cAMP significantly lower than those measured in cells expressing the MAS alleles: F142S, R231C, and L266T (Fig. 1b).

The ability of the three newly identified intragenic suppressors of R201H to support signaling through GPCRs was measured by cotransfecting cells with 2 μg of plasmid encoding the human luteinizing hormone receptor (LHR) and 5 μg of plasmid encoding one of the different isoforms of Gs, or empty vector. Protein expression was measured by western blot (Fig. 2a). Basal and hormone-stimulated levels of cAMP were measured for saturating concentrations of the LHR agonist hCG and reported as a percentage of the forskolin response (Fig. 2b). Cells expressing WT Gs exhibited a slight elevation in basal cAMP relative to cells transfected with the pCDNA3.1+ vector, and the acute response to high levels of hCG was unchanged. Expression of Gs-R201H produced a very high basal cAMP level that was not increased with hCG treatment. Other studies in our laboratory have demonstrated that transfection with lower concentrations of

Table 1  EC50 (μM) and maximum responses (% forskolin) for Gs alleles carrying R201H and intragenic suppressor mutations R231C, F142S, or L266T

<table>
<thead>
<tr>
<th>Vector</th>
<th>WT</th>
<th>R201H + R231C</th>
<th>R201H + F142S*</th>
<th>R201H + L266T</th>
</tr>
</thead>
<tbody>
<tr>
<td>EC50 (μM)</td>
<td>0.0526</td>
<td>0.179</td>
<td>0.878</td>
<td>0.00867</td>
</tr>
<tr>
<td>Emax (% forskolin response)</td>
<td>51.80</td>
<td>56.83</td>
<td>22.27*</td>
<td>51.80</td>
</tr>
</tbody>
</table>

Significantly different from WT, P < 0.05, Student’s t-test.
*Linear dose–response relationship precluded calculation of EC50 or Emax for this mutant.
Gs-R201H plasmid can produce an intermediate increase in basal cAMP that can be further increased with activation of the LHR by hCG (Tobar-Rubin et al. 2013). All three intragenic suppressors partially suppressed the response to hCG, with significant, about 50%, reductions in cAMP relative to either WT-transfected or vector-transfected cells. Dose–response curves relating cAMP production to varying doses of hCG (Fig. 2c) showed that the EC50 for hCG was 0.05 U/ml for vector-transfected cells and 0.18 U/ml for WT-transfected cells, a difference that was not statistically significant when the averages of all experiments were analyzed together (Table 1 and Supplementary Table 2, see section on supplementary data given at the end of this article). The EC50 increased to 0.88 U/ml for R201H/R231C and decreased to 0.01 U/ml for R201H/L266T. Only R201H/L266T exhibited a statistically significant change in the EC50 value (α=0.05). The maximum response to hormone was unchanged for R201H/L266T, but significantly decreased for R201H/R231C (α=0.05). R201H/F142S-transfected cells showed a linear response to hCG, precluding the calculation of EC50 or Emax.

The requirements for suppression of the R201H mutation at each of these three sites of Gs were further explored by mutating the three different suppressors to a variety of other amino acids at each site. First, F142 was mutated to alanine, tyrosine, cysteine, and aspartic acid. Cells were transfected with Gs constructs carrying each of these substitutions at position F142 along with the R201H mutation. Protein expression was confirmed by western blot (Fig. 3a), and basal cAMP levels were measured as before. All of the substitutions examined strongly suppressed the R201H mutation to 3% of forskolin levels, which was even stronger than the original F142S mutation (Fig. 3b).

For R231, the position was mutated to serine, alanine, isoleucine, glutamic acid, and lysine. Protein expression was again confirmed by western blot (Fig. 4a) and basal cAMP levels were measured (Fig. 4b). cAMP levels were not higher compared with WT (8.2±3.5% of forskolin) for R231A (15.6±6.2%), R231E (1.0±0.1%), and R231I (9.0±0.9%), while R231S (32.9±11.7%) and R231K (36.6±7.9%)
exhibited slightly elevated basal cAMP compared with WT, although still significantly lower than the original Gs-R201H mutant. Some of the low cAMP for the R231E mutation may be attributed to its lower expression levels.

Finally, mutagenesis of L266 resulted in several intragenic suppressors that were even more effective than the originally isolated mutant L266T (Fig. 5b). Substitution to aspartic acid (1.3±0.4% of forskolin), asparagine (7.5±0.4%), or serine (6.7±2.8%) resulted in basal cAMP levels not different from WT (5.3±1.9%). The conservative L266I mutation had no effect on suppressing the R201H mutation, producing basal cAMP levels of 39.3±5.7% of forskolin, not different from the R201H alone at 49.7±14.4%. Protein expression was confirmed by western blot (Fig. 5a), and the lower expression of L266D may contribute to its low basal cAMP levels. If a drug was designed to mimic the effects of these intragenic suppressors on the R201H mutation, it would also bind to the WT Gs expressed in the affected tissues. Thus, the effects of these intragenic suppressors in the absence of the R201H mutation are important to assess in order to wisely choose potential drug targets. For example, a pilot screen from our laboratory identified a suppressor that was constitutively active on its own (Tobar-Rubin et al. 2013), making that site a poor choice for drug targeting. Basal and hCG-stimulated cAMP production was measured in transfected cells expressing LHR and various mutant forms of Gs. Protein expression was confirmed by western blot (Fig. 6a). Suppressor alleles that exhibited basal cAMP levels closest to WT-transfected cells in previous experiments were chosen for these experiments. Gs isoforms carrying F142S, R231C, or L266N mutations again produced low levels of basal cAMP, and all were statistically indistinguishable from cells expressing WT Gs in their responses to maximal doses of hCG (Fig. 6b). The very low basal levels of cAMP in cells transfected with the mutant proteins were not due to inhibition of forskolin-stimulated cAMP, as the mass (pmol/well) of cAMP produced in the cells in response to forskolin was not different from the mass produced in cells transfected with WT Gs (P>0.2, ANOVA). Dose–response relationships for cAMP production in the presence of varying concentrations of hCG were determined (Fig. 6c). The EC$_{50}$ for hCG-stimulated cAMP production in all experiments combined was statistically unchanged from WT- or vector-transfected cells (Table 2 and Supplementary Table 3, see section on supplementary data given at the end of this article). R231C produced a significantly lower maximal response to hormone, while the others were unchanged from WT.

### Discussion

A yeast mutagenesis system successfully identified three completely different amino acid substitutions that are each able to suppress the constitutive activity of the disease-associated R201H mutation in Gs. These results reinforce the usefulness of yeast expression systems for large-scale random mutagenesis when asking questions relating the structure of a protein to its function (Apanovitch et al. 1998, Tobar-Rubin et al. 2013).

The three substitutions described in this paper are found in unique regions of Gs. R231, located on the Switch II helix, interacts via a salt bridge with E268, the Switch II backbone, and water molecules in the region of the GTP ligand’s phosphates (Fig. 7A and D). This mutation’s location is most similar to the location of our previously described intragenic suppressors, F222P/D223V (Tobar-Rubin et al. 2013), which was also near the GTP-binding site and interacting with water molecules associated with GTP. Mutation of R231 has been observed in patients with pseudohypoparathyroidism type 1a (Farfel et al. 1996, 2013).
Ishikawa et al. (2001), and a mild impairment of cAMP accumulation was observed in HEK293 cells transiently transfected with an R231H allele of Gs (Farfel et al. 1996). In a follow-up study, the R231H mutation was shown to destabilize GTP binding, notably blocking activation of cAMP in response to cholera toxin (Iiri et al. 1997). These studies observed that R231H proteins were ADP-ribosylated by cholera toxin, but were then unable to be activated by guanosine-5′-O-(3-thio)triphosphate (GTPγS). The activation defect was also observed in the presence of activated receptor and AlF4−, and our results add a fourth activation mechanism that is blocked by R231I mutation, mutation of R201L. The lower maximum response to hormone exhibited by R231I mutants in our studies is consistent with these previous reports.

The other two suppressor mutations are hydrophobic residues that participate in hydrophobic interactions near Switch III (L266, Fig. 7B and E) or in the helical domain of the α subunit (F142, Fig. 7C and F). Loss of hydrophobic packing may destabilize the protein, exposing it to quality control by proteolysis. However, these protein isoforms are both expressed at levels equivalent to WT and R201H forms, suggesting that protein instability is not the primary source of suppression.

The mutations at F142 are particularly intriguing. Although less studied than the Ras-like domain, the helical domain has more recently been shown to participate in G protein signaling as well (Dohlman & Jones 2012). The helical domain of Gi stabilizes nucleotide binding in the absence of Mg2+ (Coleman & Sprang 1998), and in Gs undergoes large movements away from the Ras-like domain in the structure of the heterotrimeric G-protein complexed with the agonist-occupied β2 adrenergic receptor (Rasmussen et al. 2011). Notably, F142 and its hydrophobic packing partners F140 and F146 (Fig. 7F) are located in the αB–αC loop (Switch IV), which is extended as the neighboring αB helix unwinds by a quarter turn in Giα1 activation. F142 in Gs is homologous to F118 in the Gizi3 structure, a residue that undergoes a major conformational shift from the interior of the loop in the inactive form to a solvent-exposed position in the GTPγS-bound form of Giα1 (Mixon et al. 1995). Mutation of other residues in this loop of Gs decreased GDP release, GTPγS binding, and GTP hydrolysis (Echeverria et al. 2000).

**Table 2** EC50 (µM) and maximum responses (% forskolin) for Gs alleles carrying only intragenic suppressor mutations R231C, F142S, or L266N. n = 3 independent experiments

<table>
<thead>
<tr>
<th></th>
<th>Vector</th>
<th>WT</th>
<th>R231C</th>
<th>F142S</th>
<th>L266N</th>
</tr>
</thead>
<tbody>
<tr>
<td>EC50</td>
<td>0.0693</td>
<td>0.0870</td>
<td>0.158</td>
<td>0.0340</td>
<td>0.0622</td>
</tr>
<tr>
<td>Emax (% forskolin response)</td>
<td>81.28</td>
<td>72.61</td>
<td>47.74*</td>
<td>69.59</td>
<td>59.67</td>
</tr>
</tbody>
</table>

*Significantly less than WT, P < 0.05, Student’s t-test.
Interestingly, the crystal structure of GTP\textsubscript{\gamma}S-bound Gs shows F142 packed rather than exposed to solvent (Sunahara et al. 1997), as does the structure of nucleotide-free Gs\textsubscript{-bg} G\textsubscript{2AR} (Rasmussen et al. 2011). Alignment of the helical domains of these two Gs structures shows almost no difference in the arrangement of the three phenylalanine residues in this region, even though the helical domain itself makes a large displacement.

Figure 7

The locations of the three suppressor mutations are modeled on the GTP\textsubscript{\gamma}S-bound structure of Gs (AZT1) using Chimera Software (University of California, San Francisco, CA, USA). (A, B and C) Structures of the entire Gs protein. Switch 1 is yellow, Switch II is pink, and Switch III is green. The nucleotide is shown as ball-and-stick; R201H and the indicated suppressor sites are shown in space-fill. The structures are rotated slightly left to right in (A, B and C) to provide better visualization of the location of each suppressor mutation. (D, E and F) Closeups of the local region of each suppressor mutation. (D) Interaction of Arg231 with other residues in the Ras-like domain. Hydrogen bonds are indicated in blue. Arg231 is shown colored by atom, extending from left to right from the Switch II helix. Arg231 forms a salt bridge with Glu268 (also shown colored by atom). Arg231 also hydrogen bonds to the backbone of Switch II and a water molecule. (E) Interaction of Leu266 with other amino acids in the Ras-like domain. Leu266 is shown in red, Leu270 in blue, and Phe312 is yellow (not all atoms were mapped on the structure for Phe312). These three residues pack with hydrophobic interactions in the core of the Ras-like domain of Gs. (F) Interactions of Phe142 with other residues in the helical domain. Phe142 is colored blue; Phe140 is colored yellow, and Phe146 is colored orange form a hydrophobic triad with Phe142. (G) Alignment of the helical domains of GTP\textsubscript{\gamma}S-bound Gs (1AZT) and nucleotide-free Gs (3SN6). The blue chain represents the GTP\textsubscript{\gamma}S-bound structure, with cyan residues showing the cluster of three phenylalanine residue side chains in the helical domain. The nucleotide-free structure is colored tan, with phenylalanine side chains in the helical domain colored yellow. The helical domain is to the left on the figure, showing little change in the conformations of the two protein states. The Ras-like domain is on the right of the figure, showing the large rotation of this domain relative to the helical domain in the two structures.
away from the Ras-like domain in the nucleotide-free conformation (Fig. 7G).

Several other missense mutations in Gs have been associated with hormone resistance in pseudohypoparathyroidism type 1A; L99P, P115L, C162R, R231H, S250R, R258W, R258A, Q170A, and A366S have all been reported as reducing Gs activity in these patients (MIM #103580). Similar to the mutations identified in our suppressor screen, these residues are scattered throughout the tertiary structure of Gs. Several of them (L99P, P115L, and C162R) are in the helical domain, such as the F142 suppressors reported in this paper, while another set is part of or near switch III (Q170A, S250R, and R258W), akin to the L266 site. Identification of these types of mutations in the far more complex human organismal context gives credence to our hypothesis that manipulation of the sites identified in this report may be beneficial in systems beyond simple cell culture models. Investigation of the effects of these identified suppressor sites on other alleles associated with MAS (R201C and R201S) and other constitutively active Gs alleles (Q227L) will be important follow-up studies.

The relatively high false-positive rate of identifying suppressor mutations in the yeast Gs subunit that did not suppress the constitutive activity of the R201H human Gs protein is somewhat surprising, particularly given the strong conservation of GTP-binding proteins throughout evolution. The lack of efficacy for many of the mutations identified in the yeast Gpa1 system may suggest that mutations that were successful in suppression of constitutively active Gs may be more broadly effective in suppression of any constitutively active α subunit or other activating mutations in Gs, a hypothesis that is currently under investigation in our laboratory as other α subunit activating mutations have been linked to a variety of human diseases (Landis et al. 1989, Clementi et al. 1990, Lyons et al. 1990, Lania et al. 2006).

Supplementary data
This is linked to the online version of the paper at http://dx.doi.org/10.1530/JME-13-0297.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

Funding
This work was supported by National Institutes of Health National Institute of Dental and Craniofacial Research (grant number 1R15DE020190-01) to R P-R.

References


Clementi E, Malgaretti N, Meldolesi J & Taramelli R 1990 A new subunit that did not suppress the constitutive activity of the R201H human Gs protein is somewhat surprising, particularly given the strong conservation of GTP-binding proteins throughout evolution. The lack of efficacy for many of the mutations identified in the yeast Gpa1 system may suggest that mutations that were successful in suppression of constitutively active Gs may be more broadly effective in suppression of any constitutively active α subunit or other activating mutations in Gs, a hypothesis that is currently under investigation in our laboratory as other α subunit activating mutations have been linked to a variety of human diseases (Landis et al. 1989, Clementi et al. 1990, Lyons et al. 1990, Lania et al. 2006).

Supplementary data
This is linked to the online version of the paper at http://dx.doi.org/10.1530/JME-13-0297.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

Funding
This work was supported by National Institutes of Health National Institute of Dental and Craniofacial Research (grant number 1R15DE020190-01) to R P-R.


Received in final form 5 March 2014
Accepted 17 March 2014