Evaluation of the Selectivity and Cysteine Dependence of Inhibitors across the Regulator of G Protein–Signaling Family

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ABSTRACT

Since their discovery more than 20 years ago, regulators of G protein–signaling (RGS) proteins have received considerable attention as potential drug targets because of their ability to modulate Gα activity. Efforts to identify small molecules capable of inhibiting the protein–protein interactions between activated Gα subunits and RGS proteins have yielded a substantial number of inhibitors, especially toward the well studied RGS4. These efforts also determined that many of these small molecules inhibit the protein–protein interactions through covalent modification of cysteine residues within the RGS domain that are located distal to the Gα-binding interface. As some of these cysteine residues are highly conserved within the RGS family, many of these inhibitors display activity toward multiple RGS family members. In this work, we sought to determine the selectivity of these small-molecule inhibitors against 12 RGS proteins, as well as against the cysteine-null mutants for 10 of these proteins. Using both biochemical and cell-based methods to assess Gα–RGS complex formation and Gα enzymatic activity, we found that several previously identified RGS4 inhibitors were active against other RGS members, such as RGS14, with comparable or greater potency. Additionally, for every compound tested, activity was dependent on the presence of cysteine residues. This work defines the selectivity of commercially available RGS inhibitors and provides insight into the RGS family members for which drug discovery efforts may be most likely to succeed.

Introduction

The regulators of G protein–signaling (RGS) family, comprising approximately 20 members, terminates Gα signal transduction through their activity as GTase accelerating proteins (GAP) for activated Gα subunits. This GAP activity is attributed to an approximately 120-amino-acid region common to all RGS proteins, aptly termed the RGS homology (RH) domain, which directly engages GTP-bound Gα. Since their discovery more than 20 years ago, RGS proteins have been considered attractive, but challenging, drug targets because of their ability to modulate signaling cascades occurring through a myriad of G protein–coupled receptors, including D2 dopamine (Rahman et al., 2003; Ghavami et al., 2004; Mao et al., 2004), μ opioid (Clark et al., 2003; Wang et al., 2009; Talbot et al., 2010b; Psifogeorgou et al., 2011), 5-HT1A serotonin (Ghavami et al., 2004; Talbot et al., 2010a; Wang et al., 2014), and AT1 angiotensin (Wang et al., 2002; Matsuzaki et al., 2011), to name a few. Because of their effects on signaling downstream of various G protein–coupled receptors, RGS proteins have been implicated in numerous disease states, ranging from cancer to disorders of the central nervous system. Several reviews have focused on disease states in which modulation of RGS function could be therapeutically advantageous (Hurst and Hooks, 2009; Roman and Traynor, 2011; Sjogren, 2011).

Because of the potentially beneficial effects of modulating RGS activity, considerable effort has been devoted to identifying inhibitors, particularly of RGS4, one of the most studied and perhaps best understood member of this protein family. Some of the earliest efforts focused on designing cyclic peptides that mimic the regions of Gαi that bind RGS4, leading to the discovery of the competitive peptide inhibitor YJ34 (Jin et al., 2004). This work further led to the discovery of 5nd, a peptide that covalently modified RGS4 cysteine residues (Roof et al., 2009), a theme that would become common among RGS4 inhibitors.

As development of peptides into bona fide drugs is challenging, subsequent efforts focused on using high-throughput biochemical screens to identify small-molecule inhibitors of RGS4, the first of which was Center for Chemical Genomics (CCG)-4986 (Roman et al., 2007). Follow-up work identified this compound also as being a covalent cysteine modifier (Kimple et al., 2007), albeit through binding to an allosteric site located distally to the Gα-binding face of RGS4 (Roman et al., 2010). In fact, four other RGS4 inhibitors—CCG-63802, CCG-63808, CCG-50014, and CCG-55919—were found to either interact with cysteine residues in this same region or exhibit cysteine-dependent inhibition.

Abbreviations: ANOVA, analysis of variance; CCG, Center for Chemical Genomics; Gα, guanine nucleotide binding protein, α subunit; GAP, GTase accelerating proteins; GoLoco, Gαi/o-Loco domain; NEM, N-ethyl maleimide; PPI, protein-protein interaction; RGS, regulator of G protein–signaling protein; RH, RGS homology domain; WT, XXX.
(Roman et al., 2009; Blazer et al., 2010, 2011). Previous work by our group identified UI-5 and UI-1590 as RGS4 inhibitors, both of which show lower potency against cys-null RGS4, pointing to a cys-dependent mechanism (Monroy et al., 2013). In addition to the biochemical screening methods that generally measure Gα-RGS binding or RGS-mediated GAP activity, a recent cell-based screen that measured RGS4’s ability to regulate M3 muscarinic receptor activity identified numerous inhibitors, including 6018993, 1777223, 1911669, 6383479, 5428579, and 1472216 (Storaska et al., 2013). Although none of these compounds showed activity against cys-null RGS4, both 1777223 and 5428579 exhibited reversible binding. The chemical structures of all small molecules described here and the references to their discoveries are shown in Table 1.

The RH domain is composed of nine α-helices arranged into two subdomains. The terminal subdomain in made up of helices 1–3, 8, and 9, and it contains both the N and C termini. The bundle subdomain contains helices 4–7, arranged in an antiparallel bundle. The RH domains tested here contain between zero (RGS6 and RGS7) and four (RGS2 and RGS4) cys residues, and a sequence alignment of the RH domains with cys residues highlighted is shown in Fig. 1. Notably, the cys residue located at position 95 (highlighted red, Fig. 1B) in RGS4 is conserved except for RGS6 and RGS7, where a Val residue is found. The cys residue at RGS4 position 148 (highlighted purple, Fig. 1B) is somewhat conserved, as it is found in members RGS1, RGS2, RGS4, RGS8, and RGS16.

As different RGS proteins are involved in different signaling pathways, and therefore different physiologic processes as well as potential disease states, we explored the RGS RH domain selectivity profile of previously identified RGS4 inhibitors and the reactivity of the cys residues located in this domain. Given the knowledge that nearly every known small-molecule inhibitor of RGS4 either covalently modifies cys residues or displays cys-dependent activity, we also tested each compound against a cys-null (cys to ala) mutant for every RGS tested.

**Materials and Methods**

**Compounds.** RGS-inhibiting compounds were obtained from commercial sources at indicated purity as follows: 6018993 and 1777223, both >95% purity (Vitas M Laboratories, Champaign, IL); 6383479, >90% purity (Enamine, Kiev, Ukraine); CCG-50014, >97% (Selleck, Houston, TX); CCG-63802, >98% (Sigma-Aldrich, St. Louis, MO); UI-5 and UI-1590, both ≥95% (Microsource, Gaylordsville, CT), 5428579 and CCG-4986, both ≥95% (Chembridge, San Diego, CA); CCG-63808, ≥97% (MedChem Express, Monmouth Junction, NJ); N-ethyl maleimide, ≥98% (MP Biomedicals, Santa Ana, CA); DACM, ≥95% (AnaSpec, Fremont, CA); 1472216, 98% (Key Organics, Bedford, MA); 1911669, ≥92% (Pharmeks, Moscow, Russia); CCG-55919, ≥90% (Maybridge Thermo Fisher, Waltham, MA).

**TABLE 1**

<table>
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<tr>
<th>Compound</th>
<th>Structure</th>
<th>Ref.</th>
<th>Compound</th>
<th>Structure</th>
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<td>DACM</td>
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Plasmid Construction. WT-RGS protein constructs were a gift from Nicola Burgess-Brown [Addgene plasmid no. 39143 (RGS1), 38812 (RGS2), 38813 (RGS7), 38805 (RGS8), 39138 (RGS10), 39139 (RGS14), 39140 (RGS16), and 39141 (RGS17), and 39142 (RGS18)]. Human RGS5 residues 52–185 and rat RGS4 residues 51–179 were cloned into pNIC-Bsa4 (Addgene no. 26103; gift from Opher Gileadi), as previously described (Savitsky et al., 2010). Cys-null mutants were obtained by purchasing G Blocks (Integrated DNA Technologies, Coralville, IA) containing RGS coding sequences with all cys mutated to Ala (GCG codon), which were then cloned into pNIC-Bsa4 (Addgene no. 26103), as previously described (Savitsky et al., 2010). All RGS constructs were designed to code for the RGS RH domains with an N-terminal TEV protease-cleavable 6X-His tag.

Protein Purification. All RGS proteins were transformed into BL21-CodonPlus(DE3)-RIPL cells, and colonies were selected and grown at 37°C in Terrific Broth until an OD600 of 2.0, induced with 1 mM IPTG and grown for 16 hours at 18°C while shaking at 275–300 rpm. Cells were pelleted, resuspended in 50 mM HEPES, 500 mM NaCl, 1 mM b-ME, 10 mM imidazole pH 8 (buffer A), lysed with lysozyme and DNAse I, and subjected to multiple freeze-thaw cycles in liquid N2. Lysate were clarified by centrifugation at 100,000 g, supernatant separated from insoluble pellet, and supernatant subjected to immobilized metal affinity chromatography (Ni Sepharose 6 Fast Flow; GE Healthcare, Chicago, IL). Fractions containing RGS protein were then treated with His-tagged TEV protease and dialyzed overnight at 4°C against buffer A to cleave 6X-His tag. Samples were again subjected to immobilized metal affinity chromatography and flow-through collected, resulting in >90%–95% purity as determined using SDS-PAGE (Supplemental Fig. S1). Rate-altered mutant hGo(1) (R178M, A326S) and hGo(1)-GST were purified as previously described (Monroy et al., 2013).

Chemical Biotinylation of RGS RH Domains. RGS proteins were biotinylated using EZ-link NHS-Biotin (Thermo Scientific, Waltham, MA) per manufacturer protocol with either 10:1, 5:1, or 3:1 molar excess biotin reagent and incubated on ice for 2 hours. Coupling reaction was quenched with the addition of 5000 molar excess glycine, and RGS proteins were dialyzed overnight against buffer A to remove excess glycine and biotin.

AlphaScreen Protein-Protein Interaction Assay. Biotinylated RGS was conjugated to streptavidin coated donor bead at 9X desired protein concentration and 135 ng/μl bead in Assay Buffer (20 mM HEPES, 100 mM NaCl, 1% BSA, 1% Lubrol, pH 8) on ice. After 30 minutes of conjugation the RGS/bead mixture was diluted to 3X desired concentration and added to 384-well plates (Corning 3824; Corning Scientific, Corning, NY) containing 3X concentration compound and incubated at RT. Separately, GST-Ga was conjugated to anti-GST acceptor beads at 90 nM protein and 135 ng/μl bead in assay buffer on ice. After 20 minutes of conjugation, a portion of the Ga protein/bead mixture was removed and diluted to 30 nM protein in assay buffer. This represents the negative control. The remaining Ga protein/bead mixture was diluted to 30 nM in assay buffer supplemented with 5 mM NaF, 5 mM MgCl2, 5 μM AlCl3, and 2.5 mM GDP. This mixture was incubated for an additional 10 minutes on ice before
being added to the appropriate wells of the assay plate. Final concentrations were 1× for RGS, 10 nM for Gαi1, and 15 ng/μl for each bead. Assay was incubated for 1 hour at room temperature before measurement on an Envision plate reader (PerkinElmer). All WT-RGS proteins were tested at 10 nM final. Cys-null RGS1 and RGS14 were tested at 31.6 and 100 nM, respectively, as these concentrations were necessary to achieve an acceptable difference between positive and negative control wells (data not shown).

**RGS-Gαi1 GAP Activity Assay (Malachite Green Phosphate Detection)**. Malachite green free phosphate detection assay was performed largely as described previously (Monroy et al., 2013). For assessment of RGS protein activity, RGS protein was diluted in half-log serial dilutions concentrations ranging from 3.16 μM to 3.16 nM. For compound dose-response experiments, RGS protein concentration was normalized to each RGS protein’s respective EC50, as determined above.

**WT-RGS Cysteine DACM Reactivity**. Proteins were exhaustively dialyzed against 50 mM HEPES, 500 mM NaCl at the indicated pH to remove β-ME, which can directly interact with DACM, and 20 μl of 2 μM RGS protein was then added to black 384-well plate (Corning 3575), followed by 20 μl of 2 μM DACM (Anaspec), and fluorescence intensity continuously monitored on Envision plate reader (PerkinElmer, Waltham, MA) for 1500 seconds with excitation and emission wavelengths of 385 and 440 nm, respectively.

**NanoBit Protein Complementation Assay**. The NanoBit complementation assay was performed as previously described, with minor modifications as described here (Bodle et al., 2017). NanoGlo Live Cell Reagent was prepared as a 5× stock and added to wells of the assay plate at 10 μl/well. Baseline was established for 30 minutes, after which vehicle or AlF4 (stock solution: 40 mM NaF, 500 μM AlCl3) was added to wells at 10 μl/well, and assay plate was read for an additional 30 minutes. A 4× stock of compound or DMSO was added to appropriate wells, and plate was then read for an additional hour. All reads on a Synergy 2 plate reader (BioTek, Winooski, VT) at 37°C.

**Data Analysis**. All data were analyzed using Prism 7 (GraphPad, San Diego, CA). For AlphaScreen and malachite green experiments, IC50 values were generated on the combined data set. The mean value of 3 independent experiments was determined, and these values from n = 3 independent experiments were combined in a single table to generate the mean ± S.D. values reported. Concentration-response curves were generated by a single fit on the combined mean ± S.D. data using log(inhibitor) versus response, variable slope (four parameter) fit. Data are normalized such that the absence of inhibitor was set to 100% in each assay. For AlphaScreen, 0% was normalized to the absence of GDP-AlF4, which is the negative control for protein-protein interaction (PPI) formation. Absence of RGS represented the negative control and the 0% normalization for malachite green GAP activity. After normalization, the top and bottom of each curve were constrained to 100% and 0%, respectively. IC50 values were calculated in Prism software, such that the IC50 is the concentration at which the curve crosses the 50% plane on the graph. Any calculated IC<sub>50</sub> value outside the range of concentrations (up to 100 μM) tested was identified as being >100 μM. The 95% confidence intervals were calculated by GraphPad Prism 7 using the asymmetric (likelihood) method on the single concentration response curve fit of the combined data set (mean ± S.D. data) described previously. Calculation of statistical significance was performed via one-way analysis of variance (ANOVA) with a Dunnett’s multiple comparisons post hoc analysis.

**Results**

**WT and Cys-Null RGS RH Domains Have GAP Activity**. To ensure that both the previously described and novel WT and cys-null RGS constructs produced active protein after purification (Supplemental Fig. S1), the GAP activity of each RGS protein for Gαi1 was assessed using the previously described malachite green colorimetric assay (Fig. 2) (Monroy et al., 2013). Briefly, this assay measures the GTPase activity of Gαi, by monitoring the cleavage of the γ phosphate group of GTP by Gαi1 (R178M, A326S). R178M mutation leads to decreased intrinsic GTPase activity of Gαi1 without effecting RGS sensitivity, making phosphate release RGS-dependent (Berman et al., 1996). A326S mutation increases the rate of GDP release after cleavage of the γ phosphate, the rate-limiting step in the Gα cycle, leading to receptor-independent

![Fig. 2. GAP activity of recombinant WT and Cys-null RH domains (A–D). Assessment of the concentration response curve of GAP activity of both WT and mutant RH domains of indicated RGS family members using malachite green colorimetric assay to measure GTP hydrolysis.](image-url)
EC50 values were observed. Only half that of WT RGS14 (33% vs. 64%), but overlapping cys-null (560 nM) that was more than 2-fold greater than WT.

We observed a difference in the maximal GAP activity observed. RGS5 WT exhibited greater potency (wild type [WT] 1100 AM) than cys-null (92% WT vs. 74% cys-null). RGS8 WT exhibited more potent GAP activity than cys-null, as the observed EC50 increased from 40 to 95 nM, although no differences were reached only 21% and 46% of the maximal observed activity reached for RGS5Cys was 32%, and for RGS2, this was observed activity as it exhibited no binding to G protein.

TABLE 2
Inhibition of WT-RGS: Gαq interaction/ by RGS inhibitors

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<tr>
<th>KIC50 μM</th>
<th>RGS 1</th>
<th>RGS 4</th>
<th>RGS 8</th>
<th>RGS 10</th>
<th>RGS 11</th>
<th>RGS 14</th>
<th>RGS 15</th>
<th>RGS 16</th>
<th>RGS 17</th>
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<tr>
<td>RGS 1</td>
<td>16 (12-21)</td>
<td>17 (13-24)</td>
<td>40 (30-57)</td>
<td>5.6 (2.6-8.9)</td>
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<td>0.008 (0.006-0.010)</td>
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*Data previously reported in Bodle et al., 2017b.
Furthermore, some differences between inhibition were observed in our paradigm and previous reports in the literature. Our results differ from the reports by Storaska et al. (2013) with respect to inhibition and selectivity of RGS7 and RGS8. Compound 1911669 and 5428579 were reported to inhibit RGS7, but not RGS8; but our results indicate that these compounds inhibit RGS8 but not RGS7. Compound 6383479 did not inhibit RGS8 or RGS16 using FCPIA (Storaska et al., 2013); however, we report inhibition of both proteins using AlphaScreen. It is possible that these differences can be attributed to differences in methods of FCPIA and AlphaScreen, such as incubation time, buffering conditions, RGS protein concentrations, or the amount of “active” RGS protein in the protein batch used for a given experiment.

The panel of compounds was also tested for activity against the cys-null mutant of each RGS (Supplemental Fig. S2; Table 3). Except for compounds CCG-55919 and CCG-50014, every compound was inactive (IC$_{50}$ > 100 μM) against cys-null mutants (values excluded from Table 3 for clarity). Only cys-null RGS14 was somewhat inhibited by CCG-55919 and CCG-50014, albeit at a drastically reduced potency (39-fold reduced potency CCG-55919, 5000-fold reduced potency CCG-50014). This result was as expected given the published reports detailing the activity of these compounds against cys-null RGS4.

**RGS14 is More Sensitive to Small Molecule Inhibition than RGS4.** One of the most striking results of this investigation is that although each small molecule investigated (NEM excluded) was discovered as inhibitor of RGS4, 8 of the 13 were selective for RGS14 RH at the 95% CI, whereas another (CCG-62808) was equipotent for RGS4 and RGS14 at the 95% CI. One of these compounds, CCG-50014, was previously identified as one of the most potent RGS4 inhibitors found to date, with an IC$_{50}$ of 30 nM (Blazer et al., 2011); however, the result of this investigation demonstrates that CCG-50014 inhibits RGS14 with an IC$_{50}$ of 8 nM and is selective for RGS14 at the 95% CI.

Several other compounds demonstrated moderate selectivity for RGS14 over other RGS proteins. For example, considering the UI series compounds, UI-5 was 10 times more potent for RGS14 over the next RGS; and UI-1590 inhibited RGS14 with submicromolar potency, whereas the next closest RGS was inhibited with low micromolar potency. In fact, of the 11 compounds that demonstrated inhibition of RGS14, only three (CCG-63808, CCG-4986, and CCG-55919) were not selective for RGS14 over all other RGS proteins. The apparent selectivity of these tested compounds for RGS14 is not a trait that is shared by fellow R12 family member RGS10, as only three compounds had calculable IC$_{50}$ values for RGS10. This holds true for the R4 family as well, as 11 of 13 compounds resulted in calculable IC$_{50}$ values against RGS4, whereas only one (CCG-50014) resulted in a calculable IC$_{50}$ for R4 family member RGS18.

**RGS GAP Activity Inhibition by Compounds.** Having established a biochemical pharmacological profile for known RGS inhibitors in a protein-protein interaction assay, we assessed whether the same trends would hold true in a secondary, functional biochemical assay. RGS6, RGS7, RGS10, and RGS18 were excluded due to minimal inhibition in AlphaScreen assay. Although RGS14 was readily inhibited by many of the compounds, it was excluded because 80% maximal activity could not be observed at up to 10 μM final protein concentration (Fig. 2C). As no cys-null mutants were sensitive to small-molecule inhibition in AlphaScreen, none were included.

Trends observed in the free phosphate detection assay were comparable to those observed in the protein-protein interaction assay (Fig. 3; Table 4). Compound 5428579 is not included in Table 4, as no RGS protein inhibition was observed in this assay. RGS1 and RGS4 were the most susceptible to inhibition in this assay, with six compounds resulting in calculable IC$_{50}$ values for each protein. Additionally, CCG-63808 and CCG-50014 resulted in overlapping IC$_{50}$ values for RGS1 and RGS4 at the 95% CI in this assay, where they were not overlapping in the AlphaScreen assay (Table 2). In general, for RGS1 and RGS4, the potencies of the compounds were comparable between the two assays.

This was not the case for the other four RGS proteins tested. CCG-50014 exhibited high nanomolar potencies for RGS5, RGS8, RGS16, and RGS17 using AlphaScreen but demonstrated reduced potencies in the free phosphate detection assay. Similar results were obtained for CCG-55919, as this compound resulted in low micromolar potencies using AlphaScreen and yet resulted in IC$_{50}$ values of greater than 100 μM in this GAP activity assay, possibly owing to higher protein concentrations (micromolar) required to measure GAP activity, whereas much lower concentrations (nanomolar) are needed to measure RGS: Gαi1 interaction using AlphaScreen. Alternatively, this could be caused by each assay using different Gα subunits, as AlphaScreen uses Gα$_i$, and the GAP assay a mutant of Gα$_{11}$ with accelerated GTP-binding activity. Additionally, this could be a phenomenon dependent on which RGS family member is being evaluated.

**RGS RH Domains Show Marked Differences in Cysteine Reactivity.** Because RGS inhibitors display little activity toward cys-null mutants, we sought to determine whether the relative potencies of these inhibitors were directly related to the intrinsic reactivity of their cysteine residues. For this assessment, we used the cys-reactive dye DACM, which has appreciable fluorescence only once it has covalently labeled cysteine residues in proteins (Fig. 4; Table 5). By treating the WT-RGS proteins with a 1:1 molar ratio of DACM, we observed drastically different reaction rates.
across the RGS family, which could be grossly categorized as those that readily react with DACM, those that are moderately reactive, and those that are not reactive. Independent of pH tested, RGS1, RGS2, RGS4, RGS16, and RGS18 were modified by DACM to a level that was appreciably greater than either buffer alone or the cys-devoid RGS6 and RGS7. An intermediate level of reactivity was observed for RGS14 at each pH tested and RGS10 at pH 7.0 and 6.5 but not at pH 7.5. Alternatively, RGS5, RGS8, and RGS17 interacted with DACM minimally. At every pH tested, RGS18, which

Fig. 3. Inhibition of WT RGS RH domain GAP activity by small molecules (A–G). Concentration-response assessment of the indicated inhibitor’s effect on WT RGS RH domain GAP activity using malachite green colorimetric assay to measure GTP hydrolysis. Data represent mean ± S.D. from n = 3 independent experiments and are summarized in Table 4.
contains two cys residues, was the fastest to react, although it did not display the highest relative intensity value. The next fastest rate observed was for RGS4 followed by RGS1, with RGS4 displaying lower $t_{1/2}$ and higher relative fluorescence at every pH tested, possibly because RGS4 has four cys residues versus three in RGS1. The next most reactive RH domain belonged to RGS16, which reacted approximately four times slower than RGS4 at pH 7.5, and 7.0. RGS2 reacted similarly to RGS16 at pH 7.5 but was approximately 2-fold slower at pH 7.0. Although the rate of reaction of RGS14’s two cys residues was slow (displaying an approximately linear rate), it was able to produce appreciable fluorescence at every pH, as was RGS10 at pH 6.5 and 7.0. Alternatively, RGS5, RGS8, and RGS17 showed only minimal interaction with DACM. RGS5 (one cys) reacted nearly half as fast as RGS8 (two cys residues), although, paradoxically, the $t_{1/2}$ values for both proteins did not decrease with increasing pH, as would be predicted. RGS10 and RGS17 both only have calculable $t_{1/2}$ values at pH 7.5 as a result of the curve shape, although the magnitude of these reactions (fluorescence intensity) is so low that they may be of limited relevance.

### TABLE 4

<table>
<thead>
<tr>
<th>RGS</th>
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</tr>
<tr>
<td>1</td>
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<tr>
<td>4</td>
<td>3.9 (2.4–6.0)</td>
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<tr>
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<td>20 (15–28)</td>
</tr>
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<td>16</td>
<td>19 (15–25)</td>
</tr>
<tr>
<td>17</td>
<td>31 (21–46)</td>
</tr>
</tbody>
</table>

Fig. 4. Reactivity of WT-RGS RH domains with DACM kinetic traces of the reactivity of 1 µM DACM with 1 µM indicated RGS protein normalized to RGS16 maximum. RH domains that showed appreciable reactivity are shown in (A–C), at pH 7.5, 7, and 6.5, respectively. RH domains that showed moderate to minimal reactivity are shown in (D–F) at pH 7.5, 7, and 6.5, respectively, with gray-boxed regions expanded in (G–I). Data represent mean ± S.D. from $n = 3$ independent experiments. Time to half-maximal reactivity ($t_{1/2}$) is summarized in Table 5.
Compounds 6383479 and CCG-63802 Inhibit RGS Proteins in Cells. Using a cell-based split NanoLuciferase (NanoLuc) system, we assessed the ability of several compounds to inhibit the interaction of Gα₁q with RGS proteins in cells. In this assay, one PPI binding partner is tagged with an 18-kD fragment of NanoLuc and the other binding partner a 1.3 kD NanoLuc peptide. Formation of PPI results in formation of a competent NanoLuc, and luminescence is recorded.

This system was used to test 6383479, CCG-4986, CCG-50014, and CCG-63802 against RGS1, RGS4, RGS8, RGS14, RGS16, and cysteine-null RGS4 with Gα₁q. These compounds were chosen based on the combined criteria of potency, known activity in cells and/or in vivo, and apparent RGS selectivity based on AlphaScreen and/or GAP activity results. Whereas CCG-4986 and CCG-50014 did demonstrate inhibition of several RGS proteins in this assay, this inhibition was not significantly different from the inhibition of the assay itself, as determined using a control PPI (Supplemental Fig. S3), such that no conclusions about the activity of these compounds in cells can be made. Compound 6383479 demonstrated a significant difference in signal means as determined via one-way ANOVA (F(7,16) = 10.37, P < 0.001). Dunnett’s post hoc multiple comparisons analysis revealed that RGS1, RGS4, RGS14, and cysteine-null RGS4 were inhibited by 6383479 to a significant degree compared with PPI control (P = 0.03, < 0.001, < 0.001, and 0.023, respectively) whereas RGS8 and RGS16 were not. The most robust inhibition was observed for RGS4 (Fig. 5A).

Compound CCG-63802 also demonstrated a significant difference in signal means as determined via one-way ANOVA (F(7,16) = 13.4, P < 0.001). Dunnett’s multiple comparisons analysis revealed that RGS1, RGS4, and RGS14 were inhibited by CCG-63802 to a significant degree compared with PPI control (P = 0.031, P < 0.001, and 0.001, respectively), whereas RGS8, RGS16, and cysteine-null RGS4 were not. For CCG-63802, the most robust inhibition was observed for RGS4, then RGS14, and then RGS1. Slight inhibition was also observed for RGS8, RGS16, and cysteine-null RGS4, although not to a statistically significant degree (Fig. 5B).

This cell-based system was also amenable to the RGS2-Gαq PPI. Of the four compounds tested, two inhibited RGS2 in cells (Fig. 5). Compounds 6383479 and CCG-63802 reduced the signal of the RGS2-Gαq interaction to roughly 35% and 19% of DMSO-treated controls, respectively, representing a statistically significant deviation from the control PPI (determined via one-way ANOVA as already described). Dunnett’s multiple comparison P values for each were as follows: compound 6383479, P = 0.001 and compound CCG-63802, P < 0.001. CCG-4986 and CCG-50014 did not result in inhibition of RGS2-Gαq that was discernable from their effects on the assay as was observed with Gα₁q (Supplemental Fig. S3, A–C).
Discussion

Here we present an investigation of a panel of known RGS4 inhibitors against a panel of 12 WT and 10 cys-null mutant RGS proteins, establishing the most comprehensive pharmacologic profile of these inhibitors for RGS proteins to date. Additionally, we investigated the intrinsic RGS property of cys reactivity to potentially elucidate why certain compounds are more potent against certain RGS proteins.

Investigation of 13 known RGS4 inhibitors resulted in the identification of at least one inhibitor for 10 of the 12 WT RGS proteins, with only RGS6 and RGS7 demonstrating a lack of inhibition. For RGS1, RGS2, RGS5, RGS10, RGS14, and RGS18, this represents the first identification of small molecules that inhibit these RGS proteins. Each of these RGS proteins have been directly implicated in specific pathologies (reviewed in Hurst and Hooks, 2009) or have been suggested to play a yet unverified role in pathologies (Evans et al., 2015). Therefore, through the identification of small-molecule inhibitors of these proteins, this work has provided tool compounds that may be used to probe the consequence of pharmacologic inhibition of these RGS proteins in their respective disease states. Perhaps the most powerful such tool will be CCG-50014, which we demonstrated inhibits RGS1, RGS5, RGS8, RGS14, RGS16, and RGS17 with sub-micromolar potency and has reported activity in mouse models in vivo in the investigation of RGS4 (Yoon et al., 2015). Additionally, our data greatly expand the RGS pharmacologic profile for these compounds and establishes a more comprehensive examination of compound/RGS selectivity.

The revelation that many of the small-molecule inhibitors tested were selective for the RH domain of RGS14 over that of RGS4 was surprising, as previous reports detailing the discovery of these compounds all demonstrated selectivity for RGS4. Perhaps one of the most surprising of these was CCG-50014, which was previously identified as one of the most potent small-molecule inhibitors of RGS proteins to date, with an IC$_{50}$ of 30 nM against RGS4 (Roman et al., 2009; Blazer et al., 2011). This compound was found to be a more potent inhibitor of RGS14, with an IC$_{50}$ of 8 nM and no overlap in IC$_{50}$ with RGS4 at the 95% CI. This, along with the result that many of the compounds were selective for RGS14, establishes the RH domain of RGS14 as highly sensitive to small molecule inhibition compared with the other RGS proteins tested, warranting further investigation as a target.

Given that the investigated compounds exhibited cys dependence in the reports characterizing their initial discoveries, it was not surprising that all the compounds exhibited cys dependence for all the RGS proteins. This finding led us to investigate whether the newly established pharmacologic profile with respect to these RGS proteins could be attributed to the number of cys in the RH domain of these RGS proteins. Aside from the complete absence of cys in the RH domain, the number of cys residues in the RH domain does not correlate to the potency of inhibition; there are several examples of this. RGS14 was often inhibited with the greatest potency, and yet it contains only two cys residues compared with RGS4, which contains four. Additionally, RGS18 contains two cys, one of which is highly conserved among RGS proteins and yet was only susceptible to inhibition by CCG-50014. Compare that with RGS17, which contains only one cys residue (also residing in this highly conserved region) and is inhibited by eight of the compounds tested. This led us to hypothesize that it is not the number of cys residues but rather distinct structural characteristics of the RGS proteins that alter the accessibility, and thus the reactivity of the cys residues, and that cys accessibility would correlate to observed potency of these compounds; however, our assessment of the accessibility of cys residues via use of the dye DACM demonstrates that this is not the case. In these experiments, neither the magnitude of fluorescence nor the rate of signal increase correlated to the newly established pharmacologic profile of inhibition of RGS proteins via these small molecules. This is likely because these small molecules are interacting with specific cys residues or a specific subset of cys residues in the RGS protein. Therefore, identification of the critical cys residues required for inhibition and assessment of those residues for cys accessibility may yield more interesting results. For example, RGS10 and RGS14 are closely related, as both reside in the R12 family and share >52% identity within the RH domain. RGS14 contains an additional cys residue near its C terminus in e9 and is readily inhibited by many of the compounds tested, whereas RGS10 lacks this residue and is inhibited by only three of the compounds tested, each with IC$_{50}$ > 7 μM. This provides some preliminary evidence that this C-terminal cys residue may be an important determinant for inhibitor specificity. An RGS14 construct in which the only cys present is the highly conserved cys residue in e4 could be a useful tool in future studies aimed at further understanding the molecular determinants of RGS inhibition.

Additionally, it is possible that some other structural features of the RGS RH domains confer the small-molecule selectivity established here, such as protein dynamics or altered pH values of critical cys residues. It should be noted that this work used the RH domains of all RGS proteins tested, and yet many RGS proteins are larger and more complex, containing additional domains that are sites of post-translational modification or are responsible for G protein–independent signaling cascades. The work here is critical in the investigations of variations within the RH domain that result in susceptibility to small-molecule inhibition among RGS proteins; however, investigation using the full-length constructs would shed light on the selectivity of these compounds in a more physiologically representative system, particularly for RGS family members that possess more complex multidomain and multifunctional properties. For example, many of the small-molecule inhibitors tested here were more selective for RGS14 over other RGS proteins; however, RGS14 contains a GA$_{12}/$Loco interaction (GoLoco) domain that acts as a guanosine nucleotide dissociation inhibitor and prevents dissociation of GDP from the Go subunit. Thus, the GoLoco motif adds another level of regulation of GA$_{12}$ signaling for RGS14 (Kimple et al., 2001; Willard et al., 2004). As such, the consequence of RGS14 RH inhibition on the net regulation of Go signaling in a physiologic system (i.e., using full-length RGS14) needs to be assessed.

Finally, we established a cell-based pharmacological profile for a subset of compounds using our previously characterized protein complementation assay (Bodle et al., 2017). The differential inhibition of RGS proteins observed suggests that this system may be used for the establishment of a more detailed cell based pharmacologic profile for both these and future RGS small-molecule inhibitors.
In conclusion, we have established a more comprehensive RGS pharmacologic profile for 13 published RGS4 inhibitors against a panel of 12 RGS proteins, which represents roughly 60% of the RGS family. Ten RGS proteins now have at least one identified small-molecule inhibitor that may be used as a tool compound to probe RGS function. The revelation that many small molecules inhibited RGS14 with greater potency than RGS4 demonstrates the need for more comprehensive examination of small-molecule promiscuity among RGS proteins in future discovery efforts. This work, combined with future structural interrogation, will be the foundation for establishment of RGS models that allow for design of RGS-specific small-molecule inhibitors.

Acknowledgments

The University of Iowa High Throughput Screening facility provided instrumentation, and Joseph P. O’Brien assisted in cloning and participated in helpful discussions.

Authorship Contributions

**Participated in research design:** Hayes, Bodle, Roman.

**Conducted experiments:** Hayes, Bodle.

**Contributed new reagents or analytical tools:** Hayes, Bodle, Roman.

**Performed data analysis:** Hayes, Bodle, Roman.

**Wrote or contributed to writing of manuscript:** Hayes, Bodle, Roman.

**References**


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Molecular Pharmacology

Supplemental Data

Evaluation of the Selectivity and Cysteine-Dependence of Inhibitors Across the Regulator of G Protein Signaling Family

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Department of Pharmaceutical Sciences and Experimental Therapeutics
University of Iowa, Iowa City, IA, 52242. (CRB, MPH, DLR)

Cancer Signaling and Experimental Therapeutics Program, Holden Comprehensive Cancer Center, University of Iowa Hospitals and Clinics, Iowa City, IA. (DLR)
Figure S1. SDS-PAGE analysis of purified RH domains. A. Wild-type RH domains purified using IMAC. Number above well corresponds to RGS family member. B. SDS-PAGE analysis of mutant RH domains, where all Cys residues have been replaced with Ala. Number above well corresponds to RGS family member.
Figure S2. Inhibition of WT and Cys-null RGS RH domain interactions with $\text{G}_{\alpha_0}$. Concentration-response assessment of previously described inhibitors’ ability to disrupt RGS: $\text{G}_{\alpha_0}$ PPI using AlphaScreen assay. Data for WT and Cys-null RH domains are summarized in Table 2 and Table 3, respectively. Data represent mean ± SD from n=3 independent experiments.
Figure S3. Inhibition of RGS RH domain: Goi1 interaction in cells. NanoBiT luminescence complementation assay in HEK293T cells expressing indicated RGS RH domain and Gαi1, with the notable exception that RGS2 was co-expressed with Gαq. Decrease in signal in response to compound treatment as a percent of vehicle treatment. 31.6 μM CCG-4986 (A) and CCG-50014 at 7.5 μM (B) and 31.6 μM (C) do not result RGS: Gα inhibition that is discernible from assay inhibition (Control PPI). Data represent mean of n=3 independent experiments ± SD.
Table S1

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Table S1. RGS Inhibitors Without Detectable Inhibition of WT-RGS: Gαₒ Interaction. Biochemical characterization of RGS inhibitors for their ability to disrupt the WT-RGS: Gαₒ PPI using AlphaScreen assay. Data represent the IC₅₀ with the 95% CI in parentheses from n=3 independent experiments. Concentration response curves are shown in Supplementary Figure S2. NC indicates that an IC₅₀ value was not calculable.