Regulator of G-Protein Signaling 14 (RGS14) Is a Selective H-Ras Effector

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Abstract

**Background:** Regulator of G-protein signaling (RGS) proteins have been well-described as accelerators of Gα-mediated GTP hydrolysis (“GTPase-accelerating proteins” or GAPs). However, RGS proteins with complex domain architectures are now known to regulate much more than Gα GTPase activity. RGS14 contains tandem Ras-binding domains that have been reported to bind to Rap- but not Ras GTPases in vitro, leading to the suggestion that RGS14 is a Rap-specific effector. However, more recent data from mammals and Drosophila imply that, in vivo, RGS14 may instead be an effector of Ras.

**Methodology/Principal Findings:** Full-length and truncated forms of purified RGS14 protein were found to bind indiscriminately *in vitro* to both Rap- and Ras-family GTPases, consistent with prior literature reports. In stark contrast, however, we found that in a cellular context RGS14 selectively binds to activated H-Ras and not to Rap isoforms. Co-transfection / co-immunoprecipitation experiments demonstrated the ability of full-length RGS14 to assemble a multiprotein complex with components of the ERK MAPK pathway in a manner dependent on activated H-Ras. Small interfering RNA-mediated knockdown of RGS14 inhibited both nerve growth factor- and basic fibroblast growth factor-mediated neuronal differentiation of PC12 cells, a process which is known to be dependent on Ras-ERK signaling.

**Conclusions/Significance:** In cells, RGS14 facilitates the formation of a selective Ras-GTP-Raf-ERK multiprotein complex to promote sustained ERK activation and regulate H-Ras-dependent neuritogenesis. This cellular function for RGS14 is similar but distinct from that recently described for its closely-related paralogue, RGS12, which shares the tandem Ras-binding domain architecture with RGS14.

Introduction

Many extracellular signaling molecules exert their cellular effects through activation of G protein-coupled receptors (GPCRs) [1–3]. GPCRs are seven transmembrane spanning proteins coupled to a membrane-associated heterotrimeric complex that is comprised of a GTP-hydrolyzing Gα subunit and a Gβγ dimeric partner [1,2]. Agonist-bound GPCRs catalyze the release of GDP, and subsequent binding of GTP, by the Gα subunit [1,2]. On binding GTP, conformational changes within the three ‘switch’ regions of Gα facilitate the release of the Gβγ dimer. Gα-GTP and Gβγ subunits regulate the activity of target effector proteins such as adenyl cyclases, phospholipase C isoforms, ion channels, and phosphodiesterases, which in turn regulate multiple downstream signaling cascades that initiate key biological processes such as development, vision, olfaction, cardiac contractility, and neurotransmission [1–3]. The intrinsic GTP hydrolysis (GTPase) activity of Gα resets the cycle by forming Gα-GDP—a nucleotide state with low affinity for effectors but high affinity for Gβγ. Reassociation of Gα-GDP with Gβγ reforms the inactive, GDP-bound heterotrimer which completes the cycle [1,2]. Thus, the duration of G-protein signaling through effectors is thought to be controlled by the lifetime of the Gα subunit in its GTP-bound form [2,4]. The lifetime of Gα-GTP is modulated by RGS (regulators of G-protein signaling) domain-containing proteins [4]. The RGS domain is a ~120 amino-acid nine-alpha helical bundle [5,6] that contacts Gα subunits and thereby dramatically accelerates GTPase activity [7,8]. Many RGS proteins catalyze
rapid GTP hydrolysis by isolated Gα subunits in vitro and attenuate or modulate GPCR-initiated signaling in vitro [4,5,8]; accordingly, RGS proteins are considered key desensitizers of heterotrimeric G-protein signaling pathways [4,8].

It has become apparent that the signature RGS domain is a modular protein fold found in multiple biological contexts [4,8]. The identification of multimodular RGS proteins has led to a new appreciation of these molecules as being more than just GAPs for Gα subunits [4,8,9]. RGS14 is an RGS protein with multiple signaling regulatory elements, as it contains an RGS domain, tandem RBDs (Ras-binding domains), and a GoLoco motif [10,11]. In addition to the RGS domain of RGS14 acting as a GAP for Gαi/o subunits [11–13], the GoLoco motif of RGS14 functions as a guanine nucleotide dissociation inhibitor (GDI) for Gζ subunits [14,15]. Beyond regulation of heterotrimeric Gz signaling, RGS14 is also reported to bind to activated monomeric G-proteins. An early yeast two-hybrid analysis of interactions between RGS14 and Ras-family GTPases reported a selective interaction between RGS14 and activated Rap1B, but not H-Ras [11]; in vitro experiments have also shown RGS14 binding in a nucleotide-dependent manner to the small GTPases Rap1 and Rap2 but not Ras [11,16–18]. Based on these results, it has been suggested that RGS14 may be a direct effector of Rap in vitro. However, subsequent to this initial identification of Rap (and not Ras) as a small GTPase binding target of RGS14, additional studies have suggested that Ras may also bind to RGS14. Kiel et al. [16] found that RGS14 binds preferentially to both activated Rap1B and activated H-Ras in vitro, and that this interaction is mediated by the first RBD of RGS14. Similarly, Formstecher et al. [19] identified Loco (the Drosophila RGS12/14 orthologue) in a screen for binding partners of activated Rap1, Rap2, and Ras1. Finally, we have recently discovered that RGS12, the mammalian parologue of RGS14, binds specifically to activated H-Ras in cells [20]. Collectively, these results suggest that RGS14 may bind to Rap and/or Ras GTPases. In addition to binding activated H-Ras, we found that RGS12 promotes a differentiated phenotype in both PC12 cells and embryonic DRG neurons by organizing a Ras, Raf, MEK, and ERK signal transduction complex [20]. The requirement for RGS12 in nerve growth factor (NGF)-induced neuritogenesis of PC12 cells and axonal growth of embryonic DRG neurons suggests that the related protein RGS14 may play a similar role in coordinating Ras-dependent signals that are required for promoting and/or maintaining cellular differentiation [20].

Our aim with these present studies was to resolve the discordant ideas regarding the monomeric G-protein selectivity of RGS14, as well as to establish a cellular role for such RGS14/monomeric G-protein interaction(s). Here, we demonstrate that full-length and truncated forms of RGS14 bind promiscuously to Rap and Ras GTPases in vitro, consistent with earlier reports. In cells, however, RGS14 selectively binds to activated H-Ras and not Rap nor most other Ras family isoforms. Additionally, RGS14 facilitates the formation of a Raf/MEK/ERK multiprotein complex that is dependent on activated H-Ras. Furthermore, small interfering RNA (siRNA)-mediated downregulation of RGS14 inhibits both NGF- and basic fibroblast growth factor (bFGF)-mediated neuritogenesis of PC12 cells, both processes known to require Ras-ERK signaling. These results suggest that RGS14 may regulate neuronal differentiation by the selective organization of a Ras-GTP-dependent Raf, MEK, and ERK signal transduction complex in vitro.

Materials and Methods

Materials

2.5S mouse NGF was from Roche (Indianapolis, IN). Human basic FGF was from Sigma (St. Louis, MO). Antibodies: Anti-β-actin (AC-74) and anti-FLAG M2 (Sigma), anti-ERK1/2 and anti-phospho-ERK1/2 (T202, Y204) (Cell Signaling Technologies; Danvers, MA), anti-HA-HRP 3F10 and anti-myc 9E10 (Roche), anti-myc-HRP 4A6 (Millipore, Billerica, MA), anti-rabbit IgG HRP and anti-mouse IgG HRP (GE Healthcare; Piscataway, NJ), and anti-Rap2 (BD Biosciences, San Jose, CA). All siRNAs were from Dharmaco (Lafayette, CO). siRNA sequences: rat RGS14 siGENOME SMARTpool (sense strand sequences: duplex 1, 5'-GUACCGGACUCUGUAAGCG-3'; duplex 2, 5'-GAAGAUCCACUGGCGGCGUACUG-3'; duplex 3, 5'-GGGAAGUGCUACGCCGUACG-3'. Rat RGS12/duplex 2) siRNA and the control ‘non-specific’ siRNA are described in [20]. Unless otherwise specified, all additional reagents were of the highest quality obtainable from Sigma or Fisher (Pittsburgh, PA).

Molecular biology

All DNA constructs were created using standard methods or obtained for these studies as described in Table S1. Site-directed mutagenesis was performed using the QuikChange system (Stratagene, La Jolla, CA). All DNA constructs were verified by DNA sequencing (Agencourt, Beverly, MA).

Cell culture and transfection

HEK293T and PC12 cells were cultured and transfected as described previously [20]. In PC12 experiments, pBabe-puro retrovirus expression vectors encoding constitutively-activated B-Raf(V600E) and H-Ras(G12V) were co-transfected with siRNA using LipofectAMINE 2000 (Invitrogen, Carlsbad, CA), essentially as described [20]. For co-transfections, 300 ng DNA and 150 pmol siRNA were used in a final volume of 1 ml, in 12-well plates. Neurite length was quantified after 3 days (B-Raf) or 4 days (H-Ras). We were unable to obtain an antibody capable of specifically detecting endogenous levels of murine RGS14, and thus unable to directly test for RGS14 protein knockdown in PC12 cells. To obviate this problem, we initially tested the specificity and efficacy of siRNA duplexes using HEK293T cells. HEK293T cells were plated in antibiotic-free DMEM at 165,000 cells per well in a total volume of 1 ml per well of a 12-well plate. The following day, cells were transfected with epitope-tagged RGS14 expression constructs: 20 ng myc-tagged rat RGS14 were transfected using FuGENE-6 (Roche) as described [20,21]. pcDNA3.1 was used to balance DNA amounts to a total of 1.5 μg per well. Five hours after transfection, medium was removed and cells were equilibrated in 1 ml OPTI-MEM-I (Invitrogen) for one hour. Subsequently, cells were transfected with siRNA duplexes using LipofectAMINE 2000, as described [20]. Five hours after siRNA transfection, the medium was changed to 2 ml of antibiotic-free medium per well.

Quantitative PCR

For qRT-PCR experiments that validated the specificity and efficacy of RNAi-mediated knockdown, PC12 were transfected with siRNA duplexes as described above. 48 hours post transfection, cells were washed once with PBS and then scraped and resuspended in 500 μl of PBS. Total RNA extraction and subsequent RT-PCR was performed exactly as previously described [22] using gene-specific primers and 6-carboxyfluorescein (FAM) and 6-carboxytetramethylrhodamine (TAMRA) dual labeled probes. Primer sequences: Rat β-actin: forward, 5'-TGCCGAGGATCAGTC-3'; reverse, 5'-CAGGAGACGGTGCTG-3'; probe 5'-FAM-CATCTCGGAACTAGC-3'-TAMRA3'-reverse, 5'-TGAGCCTCAG-3'; probe 5'-FAM-AAAAATAGCAGGGCCCTG-3'.
Ct(RGS14) was determined using an ABI Prism 7700 Sequence Detector. The change in the number of cycles until threshold \(C_t\) for \(RGS14\) was determined using the following equation:

\[
[\text{RGS14}] = 100 \times 2^{-[(A_0(RGS14\text{siRNA})-A_0(NS\text{siRNA}))]/(A_0(RGS14\text{siRNA})-A_0(NS\text{siRNA}))}
\]

where \(A_0(RGS14\text{siRNA}) = C_t(RGS14\text{siRNA}) - C_t(\beta\text{-actin})\) and \(A_0(NS\text{siRNA}) = C_t(NS\text{siRNA}) - C_t(\beta\text{-actin})\). All experiments were performed in triplicate.

Neurite outgrowth

PC12 neurite outgrowth was quantified essentially as described [20]. For co-transfection experiments involving siRNA knockdown along with activated H-Ras/B-Raf expression, percentages of cells containing neurites longer than one cell body were also determined. Bright-field photomicrographs of PC12 cells were obtained as described [20]. To enhance the visibility of neurites, micrographs were processed for publication using Adobe Photoshop (v7.0.1); the following commands were used sequentially: greyscale, autocontrast, autolevel, curves (50% input, 25% output).

Bimolecular fluorescence complementation

HEK 293T cells were seeded at 200,000 cells per well in 6-well dish. Cells were transfected with a total of 1 µg of DNA using FuGENE-6 (3 µl/µg of DNA). Empty pcDNA3.1 vector DNA was used to maintain a constant amount of total DNA per well. Forty-eight hours post-transfection, epifluorescence images were acquired using an Olympus IX70 fluorescence microscope with a Q-Fire CCD camera (Olympus, USA). All digital images were acquired using 1.41 sec exposures at 20× magnification and imported into Photoshop. Digital images were saved as “portable network graphics” (PNG) files and imported into MATLAB 2007a (The MathWorks, Inc. Natick, MA). Pixels with greater than 40 units of intensity in the green channel were considered to be fluorescent. The percent of fluorescent pixels for each experiment was then quantified. All experiments were repeated three times. Control experiments were performed to demonstrate the specificity of fluorescence complementation: e.g., YFP\(_N\) alone was unable to complement YFP\(_C\)-RGS14 and YFP\(_C\) alone was unable to complement YFP\(_N\)-H-Ras(G12S).

Western blotting

Protein/cell lysate electrophoresis and immunoblotting was performed as described [20]. Images were scanned using a Perfection 1200/GT-7600 scanner (Epson; Long Beach, CA). Quantification of immunoblots was performed using the Scion Image measure function (Scion Corp, Frederick, MD).

Immunoprecipitation

Immunoprecipitation experiments were conducted essentially as described [20], with the minor modification that all lysis and wash buffers contained 20 mM MgCl\(_2\). Immunoprecipitations were carried out by incubation of cell lysates with antibodies overnight at 4°C, and immune complex precipitation was achieved by incubation with 40 µl of protein A/G agarose (Santa Cruz) for one hour before washing and elution. All washing and elution steps were performed chromatographically using micro Bio-Spin columns (BioRad, Hercules, CA), as described [24]. For some experiments, pre-clearing of lysates was used to reduce non-specific binding. Pre-clearing was performed by incubating lysates at 4°C with 50 µl protein A/G agarose for 2 h. Agarose beads were removed from lysates using micro Bio-Spin columns.

GST co-precipitations

Glutathione agarose was prepared by resuspension of dry beads in excess lysis buffer (20 mM TRIS/HCL pH 7.5, 100 mM NaCl, 20 mM MgCl\(_2\), 1 mM EGTA, 1% (v/v) Triton X-100, and Complete Mini protease inhibitors (Roche)). Beads were swollen for 10 min and then washed three times by brief centrifugation, and prepared for use as a 50% (v/v) slurry. HEK293T cells were transfected with expression plasmids for small GTPases with 200 pmol GST-fusion protein was added per lysate sample, and aliquots of this mixture were taken as ‘Loading Control’ samples for SDS-PAGE. Lysate/ GST-fusion protein mixtures were then incubated overnight at 4°C, with gentle agitation. Subsequently, GST-fusion proteins and bound GTPases were precipitated with 40 µl of glutathione agarose by incubation at 4°C, for 1 h, with gentle agitation. Beads were applied to micro Bio-Spin columns and washed by gravity flow with 4×1 ml lysis buffer, followed by a final brief centrifugation (16,300×g, 30 s). Protein was eluted with 60 µl Laemmli buffer and centrifugation (16,300×g, 30 s).

Protein purification

The bacterial expression vectors pNIC-SGC/RGS14 (RBD1.RBD2) or pPROEXHTb(H-Ras) were separately expressed in E. coli, essentially as described [25]. One liter cultures of terrific broth were grown at 37°C until an OD\(_{600}\) of 1 was reached. Protein was induced with 0.5 mM isopropyl β-D-thiogalactoside for 12 h at 22°C. Cells were harvested by centrifugation at 9000×g for 20 min and resuspended in lysis buffer (50 mM HEPES, pH 7.5, 300 mM NaCl, 5% (v/v) glycerol and 10 mM imidazole) and frozen at −80°C until further use. Cell pellets containing H-Ras were resuspended in the above buffer supplemented with 2 mM MgCl\(_2\). Frozen cell pellets were thawed in the presence of one EDTA-free Complete™ protease inhibitor tablet per liter (Roche) and then were lysed using an EmulSillex C5 high pressure homogenizer (Avestin; Ottawa, Canada). Poly(ethyleneimine) was then added to a final concentration of 0.15% (v/v) and insoluble debris was removed by centrifugation for 45 min at 15000 rpm using a JA-17 rotor (Beckman Coulter, Fullerton, CA). Protein was extracted from clarified supernatant by affinity-tag purification using NTA (Ni\(^{2+}\)-nitrilotriacetate) resin (Qiagen, Valencia, California). H-Ras purification buffers were supplemented with 2 mM MgCl\(_2\). Supernatant was passed over Ni-NTA resin, which was then washed with 30 column volumes of lysis buffer and 5 column volumes of wash buffer (50 mM HEPES, pH 7.3, 300 mM NaCl, 5% (v/v) glycerol and 25 mM imidazole). Protein was eluted from the resin with 5 column volumes of elution buffer (50 mM
HEPES, pH 7.5, 300 mM NaCl, 5% (v/v) glycerol and 250 mM imidazole). Eluted protein was purified further by gel filtration chromatography using a Sephadex S200 16/60 column (GE Healthcare). RGS14(RBD1.RBD2) was subject to gel filtration using 50 mM HEPES, pH 7.5, 300 mM NaCl, and 0.5 mM Tris(2-carboxyethyl)phosphine hydrochloride. H-Ras was treated for 12 h with 50 U of calf intestinal phosphatase, 10 mM EDTA, TEV protease and 1 mM GPPNHP at 4°C and then subjected to gel filtration using 50 mM HEPES pH 7.5, 150 mM NaCl, 2 mM MgCl2, 0.5 mM TCEP. Proteins were concentrated using 10 kDa cut-off Amicon ultra filters (Millipore, Burlington, MA). GST-RGS14-His6 was purified as described [26]; all other GST-fusion proteins were purified as described [24].

Isothermal titration calorimetry

Isothermal titration calorimetry (ITC) measurements were carried out at 20°C using a VP-ITC MicroCalorimeter (MicroCal; Northampton, MA). Guanine nucleotide-loaded H-Ras and RGS14(RBD1.RBD2) were each in a solution of 20 mM HEPES, pH 7.5, 150 mM NaCl, 1 mM MgCl2 and 0.5 mM TCEP, which was degassed in a ThermoVac apparatus (MicroCal). ITC experiments were performed by stepwise titration of RGS14(RBD1.RBD2) (300 μM) into an adiabatic cell containing H-Ras (20 μM), and the heat energy change accompanying the reaction was detected upon each injection by comparison with a reference cell. Protein solution was placed in the 1.4 ml calorimeter cell and stirred to ensure rapid mixing, and 10 μl aliquots of the titrant were injected over 10 s with a 4 min interval between each injection until saturation. The titrant injected into buffer alone was used as a negative control. Heat change data was determined by subtracting values obtained when RGS14 was titrated into buffer alone. Subsequently, data was integrated and plotted against the molar ratio of H-Ras/RGS14 and analyzed as a non-linear least-squares fit. Data were analyzed using a single binding site model with the ORIGIN software package supplied by MicroCal.

Yeast two-hybrid assay

Yeast two-hybrid assays were performed essentially as described [27] using P69-4A S. cerevisiae [28]. “Bait” constructs in pGBT9 encoded the H-Ras and Rap1B GTPases fused to the Gal4p DNA binding domain, as described [29,30]. “Prey” constructs in pACTII [31] encoded the H-Ras and Rap1B GTPases fused to the Gal4p DNA binding domain, as described [29,30]. “Bait” constructs in pGBT9 encoded the H-Ras and Rap1B GTPases fused to the Gal4p DNA binding domain, as described [29,30]. “Prey” constructs in pACTII [31] encoded the Gal4p activation domain fused to the isolated RBDs of human Raf-1 or rat RGS14.

Statistics

Graphical and statistical analysis was performed using Prism 4.0 (GraphPad, San Diego, CA). All data presented are representative of three or more independent experiments.

Results

RGS14 binds promiscuously in vitro to Ras and Rap isoforms

RGS14 contains two putative RBDs in tandem, and has previously been demonstrated to interact preferentially with the GTP-bound forms of Rap1 and Rap2 but not Ras [11,17,18]. However, one group has used ITC to show that the isolated tandem RBDs of RGS14 have micromolar binding affinities for both recombinant H-Ras and Rap1B [16]. To determine the selectivity of RGS14 for Ras-family GTPases in vitro and to examine the contribution of each individual RBD to this interaction, we expressed the wild-type and activated forms of H-Ras, Rap2A, and Rap2B in HEK 293T cells, and measured the RGS14/GTPase interaction using GST pull-down assays. Purified recombinant RGS14 (both full-length and truncated versions) interacted selectively with activated (and not wild-type) H-Ras, and this interaction was dependent upon the presence of the first RBD of RGS14 (Figure 1A; e.g., compare GST-RGS14.RBD1 vs GST-RGS14.RBD2). We next examined the ability of GST-RGS14 fusion proteins to interact with wild-type and activated Rap2A and Rap2B. Interactions were observed with both Rap2A and Rap2B, and this binding appeared to be mediated by the first RBD in the tandem array; however, in contrast to the interaction with H-Ras (Figure 1A), the interaction was independent of the nucleotide state of Rap2A/2B (Figure 1B and 1C). (Note that endogenous RapGEF activity in HEK 293T cells could result in a significant amount of wild type Rap protein being GTP-bound.)

As other Ras family members can interact with RBD-containing proteins [32,33], we conducted a broader analysis of RGS14 selectivity for Ras family GTPases, initially in this in vitro setting with recombinant RGS14 protein. GST-RGS14(RBD1.RBD2) fusion protein interacted with activated versions of other Ras isoforms (K- and N-Ras) and R-Ras proteins (R-Ras1 and R-Ras3/M-Ras) in GST pull-down assays, suggesting that RGS14 is also capable of binding multiple Ras and R-Ras isoforms in vitro (Figure 2). Similarly, we examined the ability of RGS14 to interact with additional Rap isoforms. GST-RGS14(RBD1.RBD2) co-precipitated with activated Rap1A and Rap1B (Figure 2).

Isothermal titration calorimetry was then employed to demonstrate a direct in vitro protein-protein interaction and to measure the affinity of GDP or GPPNHP loaded H-Ras for RGS14(RBD1.RBD2). H-Ras was observed to directly interact with RGS14(RBD1.RBD2) in a 1-to-1 stoichiometry (Table 1 and Figure S1). The affinity between GTP-analogue bound (activated) H-Ras and RGS14(RBD1.RBD2) was significantly higher (K_D ~10 μM) than that of GDP-bound (inactive) H-Ras and RGS14(RBD1.RBD2) (K_D>200 μM) (Table 1).

RGS14 preferentially interacts with activated H-Ras in cells

We examined the capacity of RGS14 to interact with Ras proteins in mammalian cell co-immunoprecipitation (co-IP) assays. Whereas the in vitro GST pull-down assays revealed promiscuous association of full-length RGS14 (and truncated forms containing the RBDs) with multiple different Ras isoforms (Figure 1 and Figure 2), in cells full-length RGS14 stably associated preferentially with activated H-Ras over other Ras isoforms (Figure 3A and Figure S2). We consistently observed cellular co-IP of full-length RGS14 with N-Ras(G12D), but it was of lower magnitude than binding to H-Ras(G12V) (Figure S2). Interestingly, we did not observe cellular co-IP between full-length RGS14 and Rap1A, Rap1B, Rap2A, nor Rap2B (Figures S3, S4, S2), suggesting that the physiological Ras protein family target for RGS14 is H-/N-Ras, and not Rap GTPases. We also did not observe an interaction between RGS14 and activated Ras, Rap1B, Arf1, Cdc42, RalA, Rhoa, Rac1, nor Rac2 using cellular co-immunoprecipitation (Figure S2).

To examine whether full-length RGS14 and activated H-Ras form a stable complex in cells, we used yellow fluorescent protein (YFP) bimolecular fluorescence complementation [34,35]. DNA encoding N-terminal (YFP_N) and C-terminal fragments (YFP_C) of YFP were cloned in-frame with target proteins. As a positive control, we first examined the ability of YFP_N-H-Ras(G12S) and YFP_C-Raf-1 to reconstitute YFP fluorescence [36]. Cellular expression of YFP_N-H-Ras(G12S) alone or YFP_C-Raf-1 alone did not produce fluorescence (Figure 4A,B,D); however, co-expression of both proteins resulted in fluorescence complementation (Figure 4C,D). Next, we expressed YFP_N-H-Ras(G12S)
alone, YFPC-RGS14 alone, or YFPN-H-Ras(G12S) and YFPC-
RGS14, and examined reconstitution of YFP. When expressed
individually, H-Ras(G12S) and full-length RGS14 did not produce
measurable fluorescence (Figure 4E,F,H); however, when co-
expressed, the fluorescence intensity was substantially increased
(Figure 4G,H), thus demonstrating that H-Ras and RGS14
interact in live cells. We performed a comprehensive panel of
positive and negative control experiments using various YFPN and
YFPC fusion proteins (Figure S3). These controls demonstrate the
high efficiency and specificity of YFP fluorescence complementa-
tion induced by interaction between H-Ras(G12S) and RGS14. It
is of note that YFPC-RGS14 complemented YFPN-H-Ras(G12S)
with better efficiency than did YFPC-Raf-1 (Figure 4 and Figure
S3), and with comparable efficiency to the constitutive heterodi-
mer of YFPC-Gb1 and YFPN-Gc2 (Figure S3).

RGS14 coordinates an activated H-Ras-dependent B-Raf/
MEK1/ERK1 complex

To investigate the interaction of RGS14 with multiple
components of the Ras-ERK MAPK signaling pathway, we co-
expressed RGS14 and activated Ras GTases with Raf kinase
isoforms A-Raf, B-Raf, or Raf-1, and examined the ability of
RGS14 to bind to Ras. Full-length RGS14 does not interact with
activated R-Ras in cells (Figure 5A and Figure S2); however, both
activated H-Ras and R-Ras interact with all three Raf isoforms
data not shown; reviewed in [37]). Activated R-Ras did not co-
immunoprecipitate with RGS14 in the absence of any of the three
Raf kinases (Figure 5A); however, in the presence of the three Raf
kinases, we observed weak interactions with R-Ras (Figure 5A)
that were comparable to the preference of R-Ras for each of the
three Raf isoforms (data not shown). In contrast, the amount of
H-Ras bound to RGS14 dramatically increased upon concomitant
expression of B-Raf and Raf-1, but not A-Raf (Figure 5A),
consistent to our previous observations of cooperative binding with
the related protein RGS12 [20]. This interaction was specific and
not an artifact of non-specific binding of the complex to beads
(Figure S4).

We also examined whether RGS14 was able to individually or
simultaneously interact with multiple ERK MAPK components in
cells. Activated H-Ras was detected in RGS14 immunoprecipi-
tates upon their co-expression (Figure 5B and also Figure 3A). In
contrast, we did not observe binary interactions between RGS14
and B-Raf, MEK1, nor ERK1, respectively (Figure 5B). However,
when RGS14 was co-expressed with activated H-Ras, B-Raf,
MEK1, and ERK1, we isolated a complex containing all five
proteins (Figure 5B).

Loss of RGS14 inhibits NGF-mediated neurite outgrowth
in PC12 cells

Stimulation of the NGF receptor, TrkA, causes terminal
differentiation, growth inhibition, and neurite formation in PC12
cells [38,39]. NGF induces rapid and sustained activation of both
Ras and ERK, and inhibition of either Ras or ERK blocks neurite
induction [40]. Thus, NGF-induced neurite formation is mediated
by Ras activation of the ERK MAPK cascade. Loss of RGS12 (a
paralogue of RGS14) leads to reduction in NGF-promoted neurite
outgrowth of PC12 cells [20]; thus, we hypothesized that RGS14
may also play an important role in neuritogenesis in PC12 cells.
To address a possible role for RGS14 in neurite formation, we employed rat RGS14 directed-siRNA to suppress endogenous RGS14 expression. A pool of four individual duplexes efficiently reduced RGS14 expression at both the protein (Figure S5A) and mRNA levels (Figure S6B). Upon their separation, all four individual oligonucleotide duplexes also were found to efficiently knockdown expression of RGS14 (Figure S5B and Figure S6B). The RGS14-directed siRNAs did not silence RGS14 expression in PC12 cells (Figure S6A), thus demonstrating the specific nature of these reagents. RNAi-induced reduction of RGS14 expression impaired NGF-mediated neurite formation when compared to cells treated with control siRNA (Figure 6A); this led to a significant reduction in the average length of NGF-promoted neurites compared to cells transfected with non-specific siRNA (Figure 6A). bFGF can reproduce the entire spectrum of PC12 cell responses known to be elicited by NGF, including neurite outgrowth [41]; thus, we also examined whether the individual oligonucleotides were capable of reducing prolonged ERK activation by NGF and bFGF. The duration of ERK activation by NGF and bFGF was shortened by RGS14 knockdown (Figure 8E–H).

Sustained activation of ERK by NGF and bFGF is reduced upon knockdown of RGS14

In PC12 cells, sustained ERK activation promotes cell differentiation, whereas a more transient duration of ERK activation promotes growth [44–46]. Specifically, NGF, acting through the TrkA receptor, induces both transient and prolonged activation of ERK, with the prolonged activation required for neuritogenesis [46,47]. To examine the effect of RGS14 knockdown on ERK activation, PC12 cells were transfected with either non-specific siRNA or a pool of four RGS14 siRNA duplexes (Figure 8A), and stimulated with NGF or bFGF. We observed a reduction in the duration of ERK activation upon RGS14 depletion when compared to cells transfected with non-specific siRNA (Figure 8A-D). Next, we examined whether the individual oligonucleotides were capable of reducing prolonged ERK activation by NGF and bFGF. The duration of ERK activation by NGF and bFGF is reduced when compared to cells transfected with non-specific siRNA (Figure 8E–H).

**Discussion**

Our present study has generated the following major findings: (i) although RGS14 interacts with a wide array of Ras and Rap isoforms in vitro, the most likely cellular target for full-length RGS14 is activated H-Ras; (ii) the binding of activated H-Ras to RGS14 facilitates assembly of a multiprotein complex with components of the ERK MAPK cascade (B-Raf, MEK1, and ERK1); (iii) loss of RGS14 expression blunts both NGF- and bFGF-promoted neurite outgrowth of PC12 cells; and (iv) duration of ERK activation by NGF and bFGF is shortened by RGS14 knockdown, suggesting a mechanistic explanation for impairment of agonist-promoted neuritogenesis seen upon RGS14 depletion. Our findings are in contrast to the original yeast two-hybrid analysis of interactions between RGS14 and Ras-family GTPases described by Traver et al. [11], in which interaction between RGS14 and activated Rap1B, but not H-Ras, was observed. It is important to note that we have independently replicated the yeast-based data of Traver et al. [11] using (as bait) the tandem RBD of RGS14 (Figure S7). This discrepancy between yeast two-hybrid and in vitro/cellular experiments highlights the importance of examining protein-protein interactions under a variety of experimental conditions.

Our demonstration that recombinant RGS14 (both full-length and truncated species) can bind promiscuously to multiple Ras and Rap-family GTPases in vitro is not surprising as the switch

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**Table 1.** Thermodynamic parameters of H-Ras binding to the tandem Ras-binding domain region of RGS14 as measured by isothermal titration calorimetry.

| Nucleotide | $N^a$ | $K_a$ (M$^{-1}$) | $\Delta H$ (kcal M$^{-1}$) | $\Delta S$ (cal M$^{-1}$ K$^{-1}$) | $\Delta G$ (kcal M$^{-1}$) |
|------------|-------|-----------------|-----------------|-----------------|-----------------|
| GPPNHP     | 1.03 ± 0.02 | 1.01 × 10$^7$ ± 7.4 × 10$^7$ | $-1.81 ± 0.04$ | 16.6 | $-2.13$ |
| GDP        | 1.05 ± 0.06 | 2.35 × 10$^6$ ± 1.9 × 10$^6$ | $-7.1 ± 0.5$ | $-4.43$ | $-7.17$ |

$^a$Thermodynamic parameters are stoichiometry ($N$), association constant ($K_a$), enthalpy ($\Delta H$), entropy ($\Delta S$), and free energy ($\Delta G$). Data are representative of 3 or more independent experiments.

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**Figure 2.** The Ras binding domains of RGS14 have promiscuous small GTPase selectivity in vitro. GST or a GST-fusion protein of RGS14/RBD1/RBD2 were incubated with lysates from HEK293T cells transfected with mutationally activated Ras-family GTPases. Protein complexes were precipitated with glutathione agarose, washed, and resolved by SDS-PAGE and Coomassie Blue staining (top panels). Data are representative of 3 or more independent experiments.
regions of Ras-family GTPases, which participate in the interactions with Ras-binding domains, are highly conserved [48]. Yet, despite reports claiming RGS14 as a putative Rap effector [11,18], we were unable to demonstrate interaction between Rap and RGS14 in a mammalian cellular environment. We are unable to explain why the yeast two-hybrid system demonstrates that Rap1B, but not H-Ras, interacts with the RGS14 RBD region (ref. [11] and Figure S7). This suggests that, although it is a powerful discovery technique, the yeast two-hybrid system should not be used in isolation to draw conclusions about in vivo protein-protein interaction specificity. Indeed, it has been estimated that over 50% of reported yeast two-hybrid interactions are false positives [49].

Traver et al. also used purified proteins and were unable to detect an interaction between H-Ras and RGS14 [11]; we are unable to explain this difference with our present work, although we note that another group has demonstrated that H-Ras can bind to RGS14 in vitro [16]. We also note that Traver et al. may have been using low-sensitivity detection methods, as they were not able to observe interaction between RGS14 and Gz11/Gz41 [11], the latter proteins being well-established, nanomolar affinity interaction partners of the RGS14 C-terminal GoLoco motif [14]. Although we did not observe an interaction between RGS14 and Rap isoforms in cells, we have not definitively ruled out that these proteins interact in vivo. It may be that post-translational modification of RGS14 or Rap directly influences Rap/RGS14 interaction or directs these proteins to a distinct subcellular locale that facilitates their subsequent interaction [50,51].

Our observations as to the cellular selectivity of RGS14 are intriguing, in that we demonstrated that RGS14 can interact with H-Ras and, to a lesser extent, with N-Ras. Despite extensive studies, the in vivo mechanisms of Ras-effector GTPase selectivity are still not fully defined [52]. One contribution to in vivo selectivity is likely differential subcellular localizations of these GTPases, arising from

Figure 3. Full-length RGS14 interacts with activated H-Ras, but not Rap2A/B, in cells. HEK293T cells were transfected with plasmids encoding full-length, myc-epitope tagged RGS14, and wild-type (WT) or activated (GV), HA-tagged H-Ras (A), untagged Rap2A (B), or FLAG-tagged Rap2B (C). Cell lysates were immunoprecipitated (IP) with anti-myc antibodies or precipitated with GST or GST-Raf-1 (as controls). Total lysates and precipitates were immunoblotted (IB) with indicated antibodies. Data are representative of 3 or more independent experiments. doi:10.1371/journal.pone.0004884.g003

Figure 4. Full-length RGS14 and H-Ras interact in live cells. HEK293T cells were transfected with the indicated combinations of plasmids, encoding the N-terminal fragment of Yellow Fluorescent Protein (YFP) fused to the N-terminus of H-Ras(G12S), and the C-terminal fragment of YFP fused to the C-terminal of full-length RGS14, respectively. 48 h after transfection cells were analyzed by epifluorescence microscopy, and fluorescence was quantified using image analysis (as described in Materials and Methods). (A–D) Experiments measuring fluorescence complementation between H-Ras and Raf-1. (E–H) Experiments measuring fluorescence complementation between H-Ras and RGS14. Scale bars represent 50 μm. Data are representative of 3 or more independent experiments. Additional control experiments are presented in Figure S3. doi:10.1371/journal.pone.0004884.g004
post-translational modifications and/or unique hypervariable linker domain sequences outside the effector domains of Ras family members. Additionally, regions beyond the RBDs of RGS14, e.g., the RGS domain and GoLoco motif, may play a role in the selectivity of RGS14 for activated H-Ras in cells.

The apparent affinity of activated GTPases for the tandem RBD region of RGS14 in vitro is weak (e.g., for H-Ras-GPPNHP, \( K_D = 10 \mu M \)); it is thus most likely that other determinants and protein partners facilitate the formation of high affinity complexes in vivo. Despite being unable to observe binary interactions between RGS14/B-Raf, RGS14/MEK1, or RGS14/ERK1, RGS14 appears to assemble a stable, multiprotein complex containing H-Ras, B-Raf, MEK1, and ERK1 when all five proteins are expressed concomitantly (Figure 5). One report has asserted that Rap2A is unable to modulate the Gz-directed GAP or GDI activities of RGS14 in vitro [17]. However, these experiments were conducted using protein concentrations of Rap2A and RGS14 that are expressed orders of magnitude below the determined \( K_D \) values [17]. Thus it remains to be determined whether Ras-family GTPase binding to RGS14 can modulate the GAP and GDI functions of this molecule.

Our present findings with RGS14, in combination with our previous work on RGS12 [20], support the notion that both RGS proteins can function to organize multiprotein complexes containing Ras/Raf/MEK/ERK; however, how these two RGS proteins achieve this function appears different. Firstly, RGS14 does not appear to bind directly to Raf, MEK, or ERK; in contrast, RGS12 binds directly to both B-Raf and MEK2 [20]. This distinction most likely arises from the unique domain architecture of RGS12, which contains two additional domains (N-terminal PDZ and PTB domains) not present in RGS14. We established that RGS12 binds to MEK2 via its PDZ domain, and B-Raf via its tandem RBDs [20]. As RGS14 also contains tandem RBDs, it is surprising that RGS14 does not bind directly to B-Raf.

Our present data suggest that RGS14 most likely assembles a MAPK multienzyme complex differently than RGS12. This highlights the possibility that RGS14 might require additional protein partners beyond the MAPK members organized in the complex. Such a requirement for additional accessory proteins would increase the complexity of possible signaling cascades that are regulated by RGS14; it is within this scenario that RGS14 may interact with and modulate Rap-mediated signaling.

Secondly, knockdown of RGS14 in PC12 cells inhibits both NGF- and bFGF-mediated neuritogenesis, whereas depletion of RGS12 selectively inhibits only NGF-promoted neuronal differentiation. This selective modulation of growth factor receptor

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**Figure 5.** Full-length RGS14 forms a multiprotein complex with ERK MAPK pathway components dependent on activated H-Ras. (A) HA-tagged, activated H-Ras(G12V) or R-Ras(G38V) was co-transfected with empty vector, full-length myc-RGS14, or with full-length myc-RGS14 and FLAG-tagged A-Raf (“A”), B-Raf (“B”), or Raf-1 (“1”) expression vectors in HEK293T cells. Cell lysates were immunoprecipitated (IP) with anti-myc antibodies. Total lysates and immunoprecipitates were immunoblotted (IB) with indicated antibodies. (B) HEK293T cells were transfected with plasmids encoding full-length myc-RGS14, HA-H-Ras(G12V), HA-B-Raf, HA-MEK1, and HA-ERK1 in various combinations as indicated. Cell lysates were immunoprecipitated (IP) with anti-myc antibodies. Total lysates and immunoprecipitates were immunoblotted (IB) with indicated antibodies. Data are representative of 3 or more independent experiments.

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RGS14 Is an H-Ras Effector
signaling may be due, at least in part, to the ability of RGS12 to bind to the NGF receptor TrkA, but not to FGFR1 [20]. While we have shown that (a) RGS12 associates with TrkA, (b) RGS12 undergoes subcellular redistribution in response to NGF stimulation, and (c) RGS12 is localized coincident with endosomal markers in cells, we presently have no evidence for any of these functions or behaviors for RGS14. In contrast, RGS14 is typically localized to the cytosol, nucleus, and perinuclear regions in interphase, and on microtubule structures during mitosis [53–55]. Thus, coordinating activated Ras and the MAPK cascade at subcellular locales distinct from RGS12 likely engenders a different set of outputs (i.e., distinct ERK phosphorylation substrates) from RGS14-dependent signaling; independent MAPK signaling dependent on RGS14 that is equally critical for an integrated, long-term phenotypic response to a growth factor like NGF would explain why RGS12 is not able to compensate for the loss of RGS14 in NGF-induced neuritogenesis in siRNA-treated PC12 cells.

It is important to note also that RGS14 has biochemical properties atypical of a classical MAPK scaffold such as RGS12, MP1, STE5, and others. We were unable to detect binary interaction of RGS14 with any MAPK pathway members other than Ras. Typical MAPK scaffolds demonstrate binary interactions with multiple MAPK components. It is possible that H-Ras binding induces a conformational change in RGS14 that facilitates binding to additional MAPK pathway members, or that interaction with MAPK members is activation-dependent. Cellular evidence for a MAPK scaffolding-like function for RGS14 is provided by the requirement of RGS14 expression for B-Raf(V600E)-induced signal transduction in PC12 cells (Figure 7). We have not yet delineated the structural determinants of multiprotein-complex formation between Ras, RGS14, and Raf. We hypothesize that this interaction is unlikely due to simultaneous binding of RGS14 and Raf to activated H-Ras. Both RBD1 of RGS14 and the sole RBD of Raf proteins represent evolutionarily conserved binding sites for the effector loops (switch regions) of activated Ras-family GTPases [16,56]. Thus, based on the current structural knowledge, simultaneous binding of a single molecule of activated H-Ras to two RBDs is highly improbable. Evidence supporting the alternative view of Ras/RGS14/Raf complex formation is illustrated in Figure 5A, as formation of this complex is Raf-isoform selective. In the absence of RGS14, H-Ras(G12V) interacts equivalently with Raf-1, A-Raf, and B-Raf (data not shown). However, in the presence of RGS14, complex formation showed distinct selectivity towards B-Raf and Raf-1, but not A-Raf, in terms of the amount of H-Ras(G12V) co-precipitated. This suggests that facile co-precipitation of Ras in binary complexes with RGS14 and with Raf is not occurring and that a Raf-isoform selective phenomenon is being observed.

In conclusion, our studies delineate a potential major difference between the physiological roles of RGS12 and RGS14. Conventional MAPK scaffold proteins execute two main functions: (i) tethering proteins together, and (ii) specifying the subcellular localization of the multiprotein complex which, in turn, guides their final output. The finding that RGS12 is localized to endocytic vesicles and acts as a conventional MAPK scaffold that regulates NGF-promoted signaling in both PC12 and DRG neurons [20] supports the notion that RGS12 and its partners are key components of ‘signaling endosomes’ that form in the axon terminal and traffic in a retrograde manner to the cell body where...
they initiate local signal transduction cascades [57]. Thus, the subcellular localization of RGS14 is distinct from that of RGS12, and this may be reflective of functional differences in the ability to modulate signal transduction, such as the ability of RGS14, but not RGS12, to modulate FGFR-mediated signal transduction. Thus, it is likely that, in vivo, RGS14 integrates signaling independent of, and with different consequences than, RGS12.

Supporting Information

Figure S1 Guanine nucleotide-state selective interaction between H-Ras and RGS14. Isothermal titration calorimetry was used to measure the interaction between H Ras and the isolated Ras-binding domains of RGS14 ("RGS14(RBD1.RBD2)"). A stepwise titration of 300 μM RGS14(RBD1.RBD2) protein into a cell containing 20 μM H Ras(GPPNHP) (A) or H-Ras(GDP) (B) was performed and the heat change accompanying RGS14 injection was detected by comparison with a reference cell. RGS14(RBD1.RBD2) injected into buffer alone was used as a negative control. Heat changes were plotted against the molar ratio of H Ras to RGS14(RBD1.RBD2) protein and analyzed using nonlinear regression (see Table 1 of the main manuscript for data analysis parameters). Data was fit by applying a one-site binding model involving exothermic reaction phases (negative enthalpy changes) with favorable free energy changes. Analysis of the data indicates that complete saturation of the binding site is not achieved. This is likely due to the high dissociation rate of the complex.

Figure 7. RGS14 knockdown inhibits activated H-Ras- and activated B-Raf-induced neurite outgrowth. PC12 cells were co-transfected with expression vectors for constitutively-activated H-Ras(G12V) (A, B) or B-Raf(V600E) (C, D) with either a non-specific (NS) siRNA duplex or one of four independent RGS14 siRNA duplexes (#1-4). Cells left untransfected are denoted control (CTRL). Seventy-two hours after transfection, neurite outgrowth was visualized by phase contrast microscopy and digital image capture. The length of >82 neurites per condition was quantified using ImageJ. Data are plotted as dot plots with the mean of each condition represented by a black line (A, C). Significance was assessed by Kruskal-Wallis test with Dunn’s post test. For both B-Raf(V600E)- and H-Ras(G12V)-stimulated neurite outgrowth, P<0.001 for NS versus #1, #2, #3, and #4. (B, D) The percentage of cells with neurites longer than one cell body length was measured for the experiments presented in panels A and C. Data are representative of 3 or more independent experiments.

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Figure S2 RGS14 selectively interacts with H-Ras and not other small GTPases in cells. HEK293T cells were transfected with plasmids encoding full-length myc-RGS14 and various mutationally-activated ("GV", "GD", or "QL"), HA-epitope tagged GTPases. Cell lysates were immunoprecipitated (IP) with anti-myc antibodies. Total lysates and precipitates were immunoblotted (IB) with indicated antibodies. "Arf GTPase" denotes the use of Arf1.

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Figure S3 Specificity of fluorescence complementation between H-Ras(G12S) and RGS14. HEK293T cells were co-transfected with cDNAs encoding the empty vector pcDNA3.1, the N-terminal (amino acids 1–159) and C-terminal (amino acids 159–239) fragments of Yellow Fluorescent Protein (YFPN and YFPC), and indicated proteins fused to YFPN and YFPC. 48 hours after transfection, cells were analyzed by epifluorescence microscopy, and fluorescence was quantified using image analysis as described in the Experimental section. (A) Transfection of the YFP vector or YFPC-fusion constructs does not result in measurable fluorescence in the absence of YFPN co-transfection. (B) YFPN alone does not complement YFPC nor YFPC-Gβ1. (C) YFPN-H-Ras(G12S) complements both RGS14-YFPC and Raf-1-YFPC but not YFPC nor YFPC-Gβ1. (D) YFPN-Gγ2 complements YFPC-Gβ1 but not YFPC, RGS14-YFPC, nor Raf-1-YFPC.

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Figure S4 Specificity of H-Ras/ RGS14/B-Raf complex formation. HEK293T cells were transfected with plasmids encoding full-length myc-RGS14, HA-B-Raf, and either wild-type or G12V HA-H-Ras. Cell lysates were immunoprecipitated (IP) with anti-myc antibodies and protein A/G agarose, as indicated. Total lysates and immunoprecipitates were immunoblotted (IB) with indicated antibodies.

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Figure S5 Specificity and efficacy of rat RGS14 siRNAs (I). (A) HEK293T cells were transfected with HA-epitope tagged RGS14 expression vector and then 6 hours later transfected with control non-specific (NS) siRNA or a pool of four RGS14 siRNAs. 24, 48, and 72 hours later, RGS14 expression level was analyzed by immunoblot (IB) with anti-HA. Samples were immunoblotted with anti-actin antibodies as a control for total protein levels. (B) HEK293T cells were transfected with myc-epitope tagged RGS14 expression vector and then 6 hours later transfected with control non-specific (NS) siRNA or four independent RGS14 siRNA duplexes (#1-#4) that constitute the siRNA SMARTpool used in panel A. 48 hours later RGS14 expression level was analyzed by immunoblot with anti-myc antibodies. Samples were immunoblotted with anti-actin antibodies as a control for total protein levels.

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Figure S6 Specificity and efficacy of rat RGS12 and RGS14 siRNAs (II). PC-12 cells were transfected with control non-specific (NS) siRNA, RGS12(D2) siRNA (duplex 2 from ref. 20), a SMARTpool (SP) of four RGS14 siRNA duplexes, or the four individual constituent siRNA duplexes (D1, D2, D3, and D4) which comprise the SMARTpool. Forty eight hours later, cells were harvested, RNA was extracted, and RGS12 (A) and RGS14 (B) expression levels were measured by quantitative real-time PCR (as performed by the Gene Expression Core of the UNC Dept. of Laboratory Medicine and Pathology, directed by Dr. Hyung-Suk Kim). RGS12 and RGS14 data were normalized for relative expression levels using the 2-(<delta><delta>CT) method with β-actin as the internal control. Data is presented as relative expression compared to non-specific (NS) siRNA treated samples. Statistical significance was determined using ANOVA with Dunnett’s multiple comparison test (* denotes P<0.05 vs NS siRNA samples).

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Table S1 DNA constructs created and obtained for use in this study.

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Author Contributions

Conceived and designed the experiments: FSW MDW AJK RJK DD CJJD WDS DPS. Performed the experiments: FSW MDW AJK MS EAO XL RJK. Analyzed the data: FSW MDW AJK DD CJJD WDS DPS. Contributed reagents/materials/analysis tools: NAS CJD RJK. Performed the experiments: FSW MDW AJK DD CJJD WDS DPS. Contributed reagents/materials/analysis tools: NAS CJJD MJD. Wrote the paper: FSW MDW CJJD WDS DPS.

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