Disease-associated extracellular loop mutations in the adhesion G protein-coupled receptor G1 (ADGRG1; GPR56) differentially regulate downstream signaling

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Mutations to the adhesion G protein-coupled receptor ADGRG1 (G1; also known as GPR56) underlie the neurological disorder bilateral frontoparietal polymicrogyria. Disease-associated mutations in G1 studied to date are believed to induce complete loss of receptor function through disruption of either receptor trafficking or signaling activity. Given that N-terminal truncation of G1 and other adhesion G protein-coupled receptors has been shown to significantly increase the receptors’ constitutive signaling, we examined two different bilateral frontoparietal polymicrogyria-inducing extracellular loop mutations (R565W and L640R) in the context of both full-length and parietal polymicrogyria-inducing extracellular loop mutations (R565W and L640R) in the context of both full-length and N-terminally truncated (ΔNT) G1. Interestingly, we found that these mutations reduced surface expression of full-length G1 but not G1-ΔNT in HEK-293 cells. Moreover, the mutations ablated receptor-mediated activation of serum response factor luciferase, a classic measure of Gα12/13-mediated signaling, but had no effect on G1-mediated signaling to nuclear factor of activated T cells (NFAT) luciferase. Given these differential signaling results, we sought to further elucidate the pathway by which G1 can activate NFAT luciferase. We found no evidence that ΔNT activation of NFAT is dependent on Gαq/11-mediated or β-arrestin-mediated signaling but rather involves liberation of Gβγ subunits and activation of calcium channels. These findings reveal that disease-associated mutations to the extracellular loops of G1 differentially alter receptor trafficking, depending on the presence of the N terminus, and differentially alter signaling to distinct downstream pathways.

Adhesion G protein-coupled receptors (aGPCRs) are a family of GPCRs comprising 33 receptors in humans. These receptors are broadly expressed in various tissues and involved in numerous aspects of normal physiology as well as pathological processes (1). Adhesion GPCRs possess extremely long extracellular N termini that are often decorated with multiple protein/protein interaction domains. An unusual feature of these receptors is their ability to autocatalytically cleave into N-terminal fragment (NTF) and C-terminal fragment (CTF) protomers. Autoproteolysis is mediated by the GPCR autoproteolysis-inducing (GAIN) domain, which is found on the receptors’ extracellular NT near the first transmembrane domain (2). Postcleavage, the NTF and CTF remain non-covalently associated for at least some period of time. Deletion or removal of the NTF has been demonstrated to strikingly enhance the signaling activity of many aGPCRs, suggesting that the NTF normally suppresses the activity of the CTF when the two protomers are in complex (3–12).

ADGRG1 (G1; also known as GPR56) has been one of the most intensely studied aGPCRs as mutations to G1 were shown more than a decade ago to underlie the human disease bilateral frontoparietal polymicrogyria (BFPP) (13). Subsequent studies have revealed G1 to be involved in many diverse physiological processes including neurodevelopment (13–16), myelination (17), tumorigenesis (18–21), pancreatic function (22), immune function (23, 24), muscle hypertrophy (25, 26), and hematopoietic stem cell maintenance (27). To date, there are more than two dozen distinct BFPP-causing mutations (28). Although most BFPP-causing missense mutations to G1 occur on the NTF, at least five disease-associated missense mutations have been found to occur on the CTF: C418W, S485P, E496K, R565W, and L640R (13, 28, 29). In terms of functional effects, the last of those mutations (L640R) was found to ablate G1-mediated activation of RhoA following stimulation with the G1-interacting protein collagen III (30).

The activation mechanisms of aGPCRs have garnered much attention in recent years (31–33). Studies by several groups have delineated a model of activation termed the cryptic agonist model wherein dissociation of the NTF from the membrane-embedded CTF unveils the agonistic properties of the remaining extracellular stalk (also termed the “stachel”) (9–11, 34).

We previously investigated this model for G1 and found that the stalk is indeed essential for some but not all signaling outputs (35). Moreover, for other aGPCRs, such as ADGRB1 (BAII), the presence of the extracellular stalk does not appear to matter at all for receptor signaling activity (35). These findings led us to posit that aGPCRs may be capable of at least two
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![Figure 1. Schematic diagrams of full-length and ΔNT versions of R565W and L640R ADGRG1 mutant receptors.](image)

distinct modes of signaling activity: stalk-dependent and stalk-independent. For G1, the stalk was found to be required for activation of serum response factor (SRF) luciferase, a traditional measure of activity for Gα12/13-coupled receptors (36). The stalk-independent activation of nuclear factor of activated T cells (NFAT) luciferase, however, was found in those studies to rely on both G protein-dependent and -independent components and has not been clearly defined in terms of the relevant signaling cascade.

In the present study, we investigated the effects of two BFPP-causing mutations, R565W and L640R, on receptor surface expression and signaling. These studies were performed on both full-length G1 and the ΔNT truncated receptor that mimics the cleaved, active receptor. Moreover, the signaling studies assessed both stalk-dependent and stalk-independent signaling activity. The results of these studies have provided new insights into the regulation of G1 signaling and the mechanisms by which these mutations cause human disease.

### Results

**BFPP-causing mutations R565W and L640R differentially affect surface expression of full-length versus ΔNT versions of ADGRG1**

We generated four mutants version of G1: full-length (FL) and ΔNT receptors harboring either the R565W or L640R mutation (Fig. 1). The ΔNT versions of G1 lack most of the N terminus (Δ amino acids 1–382) up to the site of predicted GAIN domain cleavage and therefore mimic the CTF of G1 that is cleaved at the GAIN domain and undergoes dissociation from the NTF. We assessed the surface trafficking of each mutant in HEK-293T cells in relation to its wild-type counterpart via a cell surface biotinylation approach. As shown in Fig. 2, A and B, we observed that the surface expression and total expression of both full-length mutants were drastically reduced in comparison with the wild-type full-length receptor. Surprisingly, however, the ΔNT mutants displayed no significant deficits in surface expression compared with the wild-type ΔNT receptor (Fig. 2, C and D). It should be noted that the major band observed in Western blots for both full-length and ΔNT G1 exhibits a molecular mass of ~40–45 kDa. The full-length receptor is efficiently cleaved in HEK-293T cells, and under the denaturing conditions of SDS-PAGE utilized here, the non-covalent associations between the NTF and CTF regions are lost. Thus, Western blotting of the full-length and ΔNT receptors results in detection of the same major band as the C-terminal antibody detects the same CTF species for both versions of the receptor.

A key difference between the full-length and ΔNT receptors is that G1-ΔNT has a fully exposed extracellular stalk, whereas the stalk of the full-length receptor is mostly hidden due to either a lack of GAIN cleavage or masking by the associated NTF. The exposed stalk of G1 has agonistic properties (10, 35) and therefore may serve as a pharmacological chaperone for the receptor, counteracting the trafficking defects conferred by the R565W and L640R mutations in a manner analogous to pharmacological chaperones for other misfolded GPCRs (37). To test the hypothesis of the stalk as a pharmacological chaperone, we generated a stalkless version of the L640R G1 mutant (SL-L640R) that associates with wild-type ΔNT, L640R-ΔNT and the wild-type stalkless receptor. These data suggest that the extracellular stalk (stachel) of G1 does not act as a pharmacological chaperone as its presence made no difference for trafficking of the L640R mutant.

**R565W and L640R mutations disrupt G1-mediated activation of SRF luciferase but not NFAT luciferase**

We next assessed the signaling activity of the mutant receptors in HEK-293T cells in two distinct gene reporter assays: SRF luciferase and NFAT luciferase. In the SRF luciferase assay (Fig. 4A), none of the mutant receptors elicited significant levels of activity. In contrast, expression of the ΔNT mutant receptors resulted in substantial activation of NFAT luciferase that was comparable with the activity induced by wild-type G1-ΔNT (Fig. 4B).

Another measure of GPCR activity is association with β-arrestins (38). We previously showed that G1-ΔNT associates robustly with β-arrestin2, whereas the full-length receptor does not (4, 35). Therefore we assessed whether ΔNT-L640R could also associate with β-arrestin2 even though it is deficient in signaling to SRF luciferase. As shown in Fig. 4, C and D, co-immunoprecipitation studies revealed that ΔNT and ΔNT-L640R associate with β-arrestin2 to a similar extent, thereby providing further evidence that the mutant receptor is capable of achieving an active conformation.

**G1 signaling to NFAT luciferase does not involve β-arrestins or Gα12/13 but does involve Gβγ and calcium channels**

Given that the R565W and L640R mutations disrupted signaling to SRF luciferase but preserved signaling to NFAT luciferase and interaction with β-arrestins, we explored whether β-arrestins might be involved in mediating G1 signaling to NFAT luciferase. Overexpression of β-arrestins typically arrests G protein-dependent signaling by GPCRs but enhances...
β-arrestin-dependent signaling activity (39), and thus we studied G1 signaling in the absence and presence of β-arrestin overexpression. As shown in Fig. 5A, overexpression of β-arrestin2 significantly impaired signaling to SRF luciferase by full-length G1 and G1-NT but had no significant effect on the ability of either version of G1 to activate NFAT luciferase (Fig. 5B). These
data suggest that β-arrestins can arrest G1 signaling to SRF luciferase but are not significantly involved in G1 signaling to NFAT luciferase.

To further explore the potential role of β-arrestins in G1 signaling, we sought to remove key phosphorylation sites from the C terminus of G1 as phosphorylation of GPCR C termini is typically required for β-arrestin association (40). As a starting point for these studies, we focused on Ser-690, which is predicted by phosphorylation motif prediction algorithms to be a GPCR kinase phosphorylation site (41) and has been identified in phosphoproteomic studies to be a highly phosphorylated residue on the G1 C terminus (42). We mutated this serine to an alanine (S690A) in both FL and NT versions of G1, but subsequent co-immunoprecipitation studies revealed that NT-S690A associated with β-arrestin2 to the same extent as wild-type G1 and NT-640R displayed signaling to NFAT luciferase comparable with their wild-type counterparts. All signaling data shown here are from at least five independent experiments (**, p < 0.01; ***, p < 0.001; ****, p < 0.0001 versus cells transfected with a mock vector; error bars represent S.E.). A representative Western blot (C) and quantified results from three independent experiments (D) demonstrate that both wild-type NT and NT-640R robustly co-immunoprecipitate with HA-βArr2. IB, immunoblotting; IP, immunoprecipitation; CT, C terminus; luc, luciferase; NS, not significant.

To shed further light on G1 signaling to NFAT luciferase and how this pathway may be mechanistically distinct from G1 signaling to SRF luciferase, we performed a set of inhibitor studies. First, we assessed whether G1 might be capable of activating the Gq/11 pathway in addition to coupling to G12/13. However, as shown in Fig. 7A, we observed that U71322, an inhibitor of phospholipase C and therefore a blocker of the Gq/11 signaling cascade, had no effect on G1 activation of NFAT luciferase. Another mechanism by which GPCRs can increase cellular calcium levels to activate NFAT luciferase is via activation of plasma membrane calcium channels. Thus, we assessed G1 signaling to NFAT in the presence of SKF96365, a relatively non-specific calcium channel inhibitor (43, 44). Treatment with SKF96365 resulted in a dramatic decrease in G1-NT signaling to NFAT luciferase for both G1-NT and G1-NT-L640R (Fig. 7B). Taken together with our previous observations that Gβγ inhibitors antagonize G1-NT-mediated signaling to NFAT luciferase (35), these findings suggest that G1-NT can activate NFAT luciferase via a pathway involving the liberation of Gβγ subunits and activation of calcium channels (Fig. 7C) and moreover demonstrate that the disease-associated mutations to the G1 extracellular loops do not impair this signaling.

Discussion

In the present study, we assessed the effects of the disease-causing mutations R565W and L640R on G1 surface expression and signaling. One important observation was that the extra-
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Figure 5. β-Arrestin2 overexpression dampens ADGRG1-mediated activation of SRF but not NFAT luciferase. A, overexpression of FLAG-tagged β-arrestin2 with full-length or ΔNT G1 resulted in significant reductions in receptor-mediated activation of SRF luciferase. B, overexpression of FLAG-β-arrestin2 full-length or ΔNT G1 had no significant effect upon ADGRG1-mediated signaling to NFAT luciferase. Results are from five independent experiments (*, p < 0.05 compared with the corresponding receptor condition without FLAG-β-arrestin2; error bars represent S.E.). luc, luciferase; NS, not significant.

Several previous reports have assessed the trafficking and signaling properties of full-length BFPP-associated G1 mutants including the R565W and L640R mutants studied here (30, 47, 48). Lin and co-workers (48) found via confocal immunofluorescence that the NTF protemors for mutants R38W (distal NT), R565W (second extracellular loop), and L640R (third extracellular loop) were sharply reduced at the cell surface. In a separate study, Piao and co-workers (47) demonstrated via a cell surface biotinylation approach that surface expression of both CTF and NTF protemors was sharply reduced for mutants R38Q, R38W, Y88C (distal NT), C91S (distal NT), C346S (GAIN domain), C349S (GAIN domain), and R565W. Mixed

cellular loop mutations reduced the surface expression of full-length G1 but not the ΔNT receptor, which suggests that the tethered NTF may interact with the extracellular loops of G1. In this scenario, the R565W and L640R mutations may corrupt the normal interaction between the NTF and extracellular loops to cause protein misfolding. It is well accepted that aGPCR NTF and CTF protemors interact via hydrophobic stalk interactions within the cleaved GAIN domain (2, 45), but there has also been speculation that there may be additional NTF/CTF interactions that do not involve the stalk (31). In the case of G1, evidence in support of this idea includes the observation that the presence of the NTF strongly suppresses signaling to NFAT luciferase by the G1 CTF even though this signaling is completely stalk-independent (35). The present study provides additional evidence for NTF/CTF interactions that go beyond stalk/GAIN binding as it is unclear how the effects of extracellular loop mutations on G1 trafficking could be dependent on the presence of the NTF unless the extracellular loops possess the capacity to interact with the NTF in some way.

In addition to the effects of the R565W and L640R mutations on receptor trafficking, we also observed that these mutations ablated G1-ΔNT-mediated signaling to SRF luciferase but not NFAT luciferase. This observation suggests that the pathways by which G1 signals to SRF versus NFAT luciferase are mechanistically distinct. Indeed, we previously reported that G1-ΔNT signaling to SRF luciferase is entirely dependent on the presence of the extracellular stalk, whereas signaling to NFAT luciferase is stalk-independent (35). This previous study also demonstrated that G1 signaling to SRF luciferase was almost entirely blocked by inhibition of Go12/13, whereas signaling to NFAT luciferase was only partially dependent on Go12/13 and dependent on liberation of Gβγ subunits (35). The present study provides additional insights into the pathways downstream of G1 as our data revealed that G1 signaling to NFAT luciferase does not involve Goq/11 or β-arrestins but does involve stimulation of calcium channels in addition to Gβγ subunit liberation. Understanding the mechanism(s) by which G1 can stimulate calcium channel activity will require further elucidation, but it is interesting to note that studies on the Drosophila aGPCR lat-1 have shown that this aGPCR robustly activates transient receptor potential family calcium channels to regulate mechanosensation, perhaps via direct receptor/channel interactions (46).

Previous studies have demonstrated that NT-truncated, constitutively active aGPCRs can robustly associate with β-arrestins (4, 6, 35), but the functional effects of aGPCR interactions with β-arrestins are largely unknown. In the present study, we found evidence that β-arrestins can arrest G protein-mediated signaling by aGPCRs as β-arrestin2 overexpression dramatically inhibited G1 activation of SRF luciferase. Interestingly, however, G1 signaling to NFAT luciferase was unaffected by β-arrestin overexpression, providing yet another mechanistic distinction between these two signaling pathways downstream of G1. We also studied the functional effects of mutating a previously described (42) G1 phosphorylation site (Ser-690). Mutation of this serine residue did not alter β-arrestin association but did increase G1-ΔNT surface expression and signaling to both SRF and NFAT luciferase. These data are important because they demonstrate that G1 signaling to both SRF and NFAT luciferase is not saturated under our assay conditions. A potentially trivial explanation for the differential effects of the R565W and L640R mutations on G1-ΔNT signaling to SRF versus NFAT luciferase would be that one of these pathways was saturated, meaning that even a miniscule amount of activity in the mutant receptors might provoke a maximal amount of signaling. However, the S690A signaling data demonstrate that neither signaling pathway is saturated under the conditions of our experiments, thereby further supporting the idea that the pathways downstream of G1 to SRF versus NFAT luciferase are mechanistically distinct.
results were obtained for the L640R mutant in this study as the L640R CTF displayed a level of surface expression comparable with that of the wild-type receptor, whereas the NTF protomer was reduced at the cell surface (47). In further studies, Piao and co-workers (30) found that the L640R mutant exhibits reduced signaling relative to WT following treatment with the G1-binding protein collagen III. Our findings in the present study are consistent with the trafficking deficits that have been reported previously for the full-length R565W and L640R mutants. However, the present study also significantly extends work in this area with the surprising observation that the deleterious effects of these mutations on G1 trafficking are completely abrogated in the ΔNT form of the receptor. Additionally, we found that, although the activated L640R mutant receptor is deficient in Gα12/13-mediated signaling as Piao and co-workers (30) observed, this receptor still robustly binds to β-arrestins and can activate NFAT luciferase via stimulation of calcium channel activity. Thus, our data suggest that the L640R mutant receptor is not completely inactive but rather selectively deficient in certain aspects of its signaling.

In summary, the present study has provided novel insights into how disease-associated mutations to the G1 extracellular loops can differentially impact G1 trafficking and signaling. Going forward, it will be of interest to study further whether aGPCR extracellular loops do indeed interact with the tethered NTF regions as suggested by the findings reported here and to understand what the structural basis of these interactions may be. Additionally, another point of interest will be to further dissect stalk-dependent versus stalk-independent modes of aGPCR signaling and to understand how aGPCR extracellular regions (the NTF and extracellular loops) can differentially modulate distinct aspects of receptor signaling. Finally, a major goal of fundamental studies into aGPCR signaling like those reported here is to set the stage for the future pharmacological targeting of aGPCRs with small molecule agonists, antagonists, and modulators. Given the importance of this family of receptors

Figure 6. Mutation of a putative phosphorylation site (S690A) on the C terminus of ADGRG1 enhances surface expression and signaling by the ΔNT mutant but does not abolish binding to β-arrestin2. A representative Western blot (A) and quantified results from three independent experiments (B) demonstrate that there was no significant difference in co-immunoprecipitation with β-arrestin2 between wild-type G1-ΔNT and ΔNT-S690A. A representative Western blot (C) and quantified results from three independent experiments (D) reveal that the S690A mutation enhanced the surface expression of the ΔNT mutant but not the full-length mutant. The ΔNT-S690A mutant also displayed significantly higher levels of SRF (E) and NFAT luciferase (F) activation compared with the wild-type ΔNT receptor (*, p < 0.05; **, p < 0.01 for the indicated comparisons; error bars represent S.E.). Results shown are from at least four independent experiments. IB, immunoblotting; IP, immunoprecipitation; CT, C terminus; NS, not significant; luc, luciferase; a.u., arbitrary units.
for human health and disease (1), the members of this family may prove to be important drug targets for novel classes of therapeutics in the treatment of many different human diseases.

**Experimental procedures**

**Constructs**

Human G1-ΔNT (amino acids 383–693) and G1-SL (amino acids 404–693) were subcloned into pcDNA3.1 between 5′ HindIII (G1-ΔNT, GCAAAGAAGCTTATGACCTACTTTG-CAGTGCTGATG; G1-SL, GCAAAGAAGCTTATGAGCCTCCTCTCCTACGTGGG) and 3′ XbaI (GCAAAGTCTAGACTAGATGCGGCTGGACGAGGT). FLAG- and HA-arrestin2 constructs have been described previously (35). R565W mutant receptors were generated using the following primers: 5′-CCATGTGCTGGATCTGGGACTCCCTGGTC-3′ and 5′-GACCAGGGAGTCCCAGATCCAGCATGG-3′. L640R mutant receptors were generated using the following primers: 5′-TGATGCTGAAAAGGTAGCGGAGACAAGCTGGAAG-3′ and 5′-CTTCCAGCTTGTCGTCGCTACCTTTCAGCATCA-3′. S690A mutant receptors were generated using the following primers: 5′-AGATGCGGCTGGCCAGGCTGCTGAGG-3′ and 5′-GGCAGCACCCTCGGCCAGCCGCAGCTCT-3′. All mutant receptors were generated using the QuikChange Lightning site-directed mutagenesis kit (Agilent, catalog number 210519) according to the manufacturer’s protocol.

**Reagents**

SKF 96365 was purchased from Cayman Chemicals (catalog number 10009312), and U73122 was purchased from Tocris Biosciences (catalog number 1268). All other general reagents were from Sigma.

**Cell culture**

HEK-293T/17 cells were acquired from ATCC (Manassas, VA) and maintained in DMEM (Life Technologies) supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin in a humid, 5% CO₂, 37°C incubator. Cells were transfected using Mirus (Madison, WI) TransIT-LT1 according to the manufacturer’s protocol.

**Western blotting**

Protein samples were reduced and denatured in Laemmli buffer, loaded into 4–20% Tris-glycine gels (Bio-Rad) for SDS-
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PAGE, and then transferred to nitrocellulose membranes (Bio-Rad). Blots were blocked with 5% milk (in 50 mM NaCl, 10 mM HEPES, pH 7.3, with 0.1% Tween 20) and incubated with primary antibodies for 1 h at room temperature or overnight at 4 °C. The anti-GPR56 C-terminal antibody was developed by Orbigen, Inc. via injection of rabbits with a peptide (CSNSD-SARLPISSGSTSSSRI) derived from the GPR56 C terminus and has been characterized previously (4). Rat anti-HA (Roche Applied Science) and mouse HRP-conjugated anti-FLAG (Sigma) antibodies were used to detect epitope-tagged proteins. HRP-conjugated secondary antibodies were purchased from GE Healthcare, and antibody labeling of specific bands was visualized using Thermo Scientific SuperSignal West solutions.

Cell surface biotinylation

HEK-293T cells were transfected with 2 μg of DNA (empty vector or receptor). At 24-h post-transfection, cells were placed on ice and washed with ice-cold PBS + Ca2+ three times. Cells were then incubated with 10 mM Sulfo-NHS-Biotin (Thermo Scientific) in PBS + Ca2+ on ice for 30 min and then washed three more times with PBS + Ca2+ + 100 mM glycine. Cells were resuspended in 250 μL of lysis buffer (1% Triton X-100, 25 mM HEPES, 150 mM NaCl, 10 mM MgCl2, 1 mM EDTA, protease inhibitor mixture (Roche Diagnostics), 2% glycerol) and lysed by slowly rotating on a spinning wheel for 30 min at 4 °C. Cell debris was cleared by centrifugation, and soluble cell lysates were incubated with 50 μL of streptavidin-agarose beads (Thermo Scientific) for 1 h at 4 °C. Beads were washed three times with lysis buffer and resuspended in 60 μL of Laemmli buffer. Biotinylated proteins were detected via Western blotting as described above. Western blot bands were quantified using Image Studio software (LI-COR Biosciences, Lincoln, NE).

β-Arr2 binding assay

HEK-293T cells were transfected with a total of 6 μg of DNA (empty vector, receptor, and/or HA-β-arrestin2 (βAr2) or FLAG-βAr2). The next day, cells were washed with cold PBS + Ca2+ and lysed in harvest buffer (150 mM NaCl, 25 mM HEPES, pH 7.3, 1 mM EDTA, 10 mM MgCl2, 1% Triton X-100, Roche EDTA-free Complete protease inhibitor mixture tablet). Lysates were rotated at 4 °C for 45 min to solubilize integral membrane proteins, and membranes were cleared by centrifugation (15 min at 17,000 × g at 4 °C). Solubilizates were added to anti-HA- (Sigma) or anti-FLAG-agarose beads (Sigma) and rotated at 4 °C for 1 h. Beads were washed three times in harvest buffer, and proteins were eluted in Laemmli buffer at 37 °C for 10–15 min and loaded in 4–20% Tris-glycine gels for SDS-PAGE and Western blotting. Western blot bands were quantified using Image Studio software.

Luciferase reporter assays

HEK-293T cells were seeded in 96-well plates 20–24 h prior to transfection. Each well was transfected with 50 ng of firefly reporter, 1 ng of Renilla luciferase, and 10 ng of receptor or empty plasmid DNA. All reporter constructs (NFAT, pGL4.30; SRF, pGL4.34; Renilla, pRLSV40) were acquired from Promega (Madison, WI). At 24–48 h later, Dual-Glo luciferase assays (Promega) were performed according to the manufacturer’s protocol, and plates were read on a BMG Omega plate reader. For inhibitor studies, U73122 (10 μM; diluted from a stock in DMSO) or SKF96365 (50 μM; diluted from a DMSO stock) were added to wells for 8 h before the plates were read. Vehicle control wells received an equivalent amount of DMSO (0.1% final). Results were calculated for each assay by determining the luminescence ratio of firefly:Renilla luciferase counts normalized to empty vector-transfected wells. Error bars for all empty vector-transfected conditions were represented as the standard errors of the normalized raw value means.

Author contributions—A. K. and R. A. H. designed experiments and wrote the manuscript. A. K. performed the experiments. A. K. and R. A. H. analyzed the data and created the figures. All authors made intellectual contributions to the paper and take responsibility for the data.

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