Alternative Translation Initiation of Human Regulators of G-Protein Signaling-2 Yields a Set of Functionally Distinct Proteins

Steven Gu, Annepa Anton, Samina Salim, Kendall J. Blumer, Carmen W. Dessauer, and Scott P. Heximer

Department of Physiology, Heart and Stroke/Richard Lewar Centre of Excellence in Cardiovascular Research, University of Toronto, Toronto, Ontario, Canada (S.G., A.A., S.P.H.); Department of Integrative Biology and Pharmacology, University of Texas Health Science Center at Houston, Houston, Texas (S.S., C.W.D.); and Department of Cell Biology and Physiology, Washington University School of Medicine, St. Louis, Missouri (K.J.B.)

Received March 22, 2007; accepted September 10, 2007

ABSTRACT

The regulator of G-protein signaling (RGS2) contains a characteristic RGS domain flanked by short amino and carboxyl terminal sequences. The RGS domain mediates inhibition of \( \alpha \)-subunit activity via GTPase activating protein (GAP) domains (Berman et al., 1996; Watson et al., 1996) and thus are important for signal modulation and discrimination. A number of RGS proteins contain activities that extend beyond their GAP function. Proteins within the RGS7-like, RGS12-like, Rhogef-containing, and G protein-coupled receptor kinase-like RGS protein subfamilies contain multiple modular protein-protein interaction domains that allow them to coordinate signaling between intracellular signaling networks (Zheng et al., 1999; Ross and Wilkie, 2000). By comparison, simply constructed RGS proteins in the RGSZ-like and RGS4-like (R4/B) subfamilies consist of little more than an RGS domain flanked by short (typically 10–70 residues) amino- and carboxyl-terminal extensions. It is evident that even such simple RGS proteins can be versatile.

Heterotrimeric G-protein-coupled receptors mediate cell responses to a variety of extracellular ligands (Ma and Zemmel, 2002). Coordination of G-protein signaling allows cells to adjust rapidly to dynamic physiological conditions. Mammalian regulators of G-protein signaling (RGS) proteins attenuate G-protein \( \alpha \) subunit activity via GTPase activating protein (GAP) domains (Berman et al., 1996; Watson et al., 1996) and thus are important for signal modulation and discrimination. A number of RGS proteins contain activities that extend beyond their GAP function. Proteins within the RGS7-like, RGS12-like, Rhogef-containing, and G protein-coupled receptor kinase-like RGS protein subfamilies contain multiple modular protein-protein interaction domains that allow them to coordinate signaling between intracellular signaling networks (Zheng et al., 1999; Ross and Wilkie, 2000). By comparison, simply constructed RGS proteins in the RGSZ-like and RGS4-like (R4/B) subfamilies consist of little more than an RGS domain flanked by short (typically 10–70 residues) amino- and carboxyl-terminal extensions. It is evident that even such simple RGS proteins can be versatile.

ABBREVIATIONS: RGS, regulator of G-protein signaling; GAP, GTPase activating protein; AC, adenylyl cyclase; ACV, type V adenylyl cyclase; NTD, amino terminal domain; NT, amino terminus; ORF, open reading frame; GFP, green fluorescent protein; ROI, region of interest; IBMX, 3-isobutyl-1-methylxanthine; TMRM, tetramethyl rhodamine methyl ester; RFU, relative fluorescent unit; AM, acetoxymethyl ester; EGFP, enhanced green fluorescent protein; wt, wild type; HEK, human embryonic kidney; FR, fluorescence ratio; IPx, inositol phosphate; kz, Kozak; MG132, N-benzyloxy carbonyl (Z)-Leu-Leu-leucinal.


The online version of this article (available at http://molpharm.aspetjournals.org) contains supplemental material.
integrators of G-protein signaling through their interaction with a diverse number of intracellular protein partners (Heximer and Blumer, 2007).

RGS2 belongs to the R4/B subfamily of simple RGS proteins. Despite its small size, RGS2 can interact with G-proteins and non–G-protein signaling partners. The GAP domain of RGS2 inhibits Goα (Heximer et al., 1999) and Goβ (Ingi et al., 1998) signaling, whereas sequences within the RGS2 amino terminal domain (NTD) direct nuclear and plasma membrane targeting (Heximer et al., 2001). More recently, however, the NTD of RGS2 has also been shown to interact with additional signaling partners including G-protein-coupled receptor third intracellular loops and spinophilin (Bernstein et al., 2004; Hague et al., 2005; Wang et al., 2005), adenylyl cyclase (Sinaramjah et al., 2001; Salim et al., 2003; Roy et al., 2006a), TRPV6 (Schoeber et al., 2005; Liao et al., 2006; Liao et al., 2007) and eotropic G(βγ) subunits (Kozak et al., 1989; Kozak, 1992). Together, these observations prompted us to study alternative mRNA splicing. Chidiac and coworkers recently showed that multiple RGS2 proteins are expressed in the NTD of RGS2 with various functional properties is expected to direct the carboxyl terminal GAP domain into context-specific signaling compartments.

Several RGS genes produce more than one protein with unique functional properties using alternative mRNA splicing. Chidiac and coworkers recently showed that multiple RGS2 bands were evident in forskolin-stimulated mouse osteoblasts (Roy et al., 2006b). We examined the possible mechanisms that might result in the production of multiple RGS2 proteins. A search of the human-expressed sequence tag database (National Library of Medicine, National Center for Biotechnology Information) revealed no evidence of alternatively spliced RGS2 mRNAs. Furthermore, our own expression data suggested that multiple RGS2 proteins are expressed from the full RGS2 cDNA alone (Heximer et al., 1999). Together, these observations prompted us to study whether alternative translation of the human RGS2 mRNA was important for the regulation of its expression and function. Here, we report the discovery of a novel set of alternatively translated RGS2 proteins with distinct functional properties whose relative expression levels are coupled to changes in cell signaling status.

Materials and Methods

Materials. The pEGFP-C1 or pREV-TRE (Clontech, Mountain View, CA) plasmids were used to express RGS2 in this study. Constitutively active Goα (R183C) construct in pCIS was a kind gift from Dr. J. Hepler (Emory University, Atlanta, GA). Constitutively active Goα (G Q227L) and the ACV clone were kindly provided by Drs. R. Feldman and P. Chidiac (University of Western Ontario, London, ON, Canada). Expression constructs for Fibrillarin-HcRed were kindly donated by Dr. K. Lukyanov (Shemyakin and Ovchinnikov Institute of Bioorganic Chemistry, Moscow, Russia). Polyclonal anti-green fluorescent protein (GFP) antibody was from Clontech, and horseradish peroxidase-coupled goat anti-rabbit IgG secondary and mouse 9E10 monoclonal anti-myc epitope antibodies were from Covance Research Products (Denver, PA). HEK293 cells stably expressing the M1 muscarinic receptor were kindly provided by P. Burgon and E. Peralta. Tet-ON HEK293 cells were from Clontech. Tet ON-HEK293 cell lines were maintained in Dulbecco's modified Eagle's medium: Ham's F-12 medium (1:1) and α-minimal essential medium, respectively, supplemented with 10% (v/v) heat-inactivated fetal calf serum (Atlanta Biologicals, Lawrenceville, GA), 2 mM glutamine, 10 µg/ml streptomycin, and 100 U/ml penicillin at 37°C in a humidified atmosphere with 5% CO₂. For doxycycline-induction studies, transiently transfected Tet-ON HEK293 cells were treated for 48 h with the indicated doxycycline concentrations before harvesting for immunoblotting. All stably transfected HEK293 cell lines expressing epitope-tagged RGS2 were generated essentially as described previously (Heximer et al., 1999). In brief, a clonal population HEK293 cells (7 × 10⁶ cells in 10-cm plates) was transfected with 5 µg of mammalian expression constructs that express direct translation of expression start-site optimized and wild-type RGS2 constructs that had been tagged at their carboxyl termini with three tandem copies of the c-myc epitope. Cells were plated at limiting dilution, and stable RGS2-(myc)-expressing clones were selected for in growth medium containing 0.5 mg/ml Geneticin. Cell lines expressing similar levels of RGS2 protein were identified by Western blotting, and clonality was verified by immunofluorescence staining using the mouse 9E10 monoclonal antibody. Clonal cell lines were immediately frozen in aliquots for storage at passage 3 to 4. The possibility of that loss of the appropriate signaling molecules occurred during clonal selection was minimized by examination of the relevant signaling readouts in 22 separate control (3 lines) and RGS (19 lines)-expressing cell lines. All vector control lines showed similar signaling efficiency. Two RGS-expressing clones showed greater inhibition than expected from their apparent low levels of expression and were not included. To determine the relative expression levels of RGS proteins in stably transfected cell lines, cells from trypanspized plates were counted, pelleted, and lysed (2 × 10⁶ cells/ml) in Laemmli sample buffer and resolved by SDS-polyacrylamide gel electrophoresis. RGS2 protein expression patterns were determined by immunoblotting using antibodies directed against the indicated epitope tag in phosphate-buffered saline (PBS)/0.1% Tween-20 (PBST)/Tween-20 (PBST) containing 3% (v/v) bovine serum albumin (anti-GFP, 1:400; anti-myc, 1:1000) and enhanced chemiluminescence. Where indicated, densi-
tometric quantitation of protein expression was performed using the gel analysis function of the Image 1.32 software package.

**Phosphoinositide Hydrolysis Assays.** Inositol phosphate accumulation in stably and transiently transfected cell lines was measured essentially as described previously (Heximer, 2004).

**Intracellular Calcium Imaging.** HEK293 cells stably transfected with the M1 muscarinic receptor were seeded at 50% confluence on polylysine-coated #1 glass coverslips in 6-well plates before transfection with 1 μg of plasmid DNA in FuGENE 6 (Roche). After 24-h transfection, coverslips were washed and incubated in calcium imaging buffer (11 mM glucose, 130 mM NaCl, 4.8 mM KCl, 1.2 mM MgCl₂, 17 mM HEPES, and 1 mM CaCl₂, pH 7.3) containing 5 μM fura-2 AM and 0.05% Pluronic acid for 40 min at 37°C. Fura-2-loaded cells were washed again and incubated for at least 10 min in calcium imaging buffer to allow hydrolysis of the AM ester. Coverslips were mounted in a TC1-2SL25 open-bath chamber (BioScience Tools, San Diego, CA) and imaged on an Olympus BX51WI upright microscope (Olympus, Tokyo, Japan) using a 10× water-dipping objective. Excitation light was provided by a DeltaRam V monochromator (PTI, Lawrenceville, NJ). Fluorescence imaging was performed with ImageMaster imaging software (PTI). Images were acquired with a Photometrics Cascade 512B cooled charge-coupled device camera (Roper Scientific, Tucson, AZ). GFP and RGS2-GFP expressing cells were identified using 488 ± 5 nm excitation and selected as regions of interest (ROIs). Relative GFP fluorescence (RGS expression) and fura-2 ratiometric (intracellular calcium) was determined for each ROI and was calculated as mean pixel fluorescence value after 200- and 100-ms exposure, respectively. For fura-2 imaging, alternating excitation wavelengths (355 ± 5/396 ± 5 nm) were provided at −1 excitation pair per second and paired images collected through a 510 ± 20-nm emission filter (Chroma Technology Corp., Brattleboro, VT). Fluorescent ratio (FR) values for the images pairs were determined for ROIs selected on the basis of their GFP expression. Baseline fluorescence ratios of nonstimulated cells were collected from 30 frames before the addition of 200 μM carbachol. The percentage of increase from baseline FR levels to the peak stimulated FR was determined specifically for low GFP or RGS2-GFP-expressing cells with relative GFP fluorescence between background levels (3300 relative fluorescent units, RFU) and an upper experimental limit of 10,000 RFU. Higher expression levels provide greater (even complete) attenuation of the intracellular calcium response; however, high intracellular GFP levels result interfere with the 396 nm channel during fura-2 excitation. For technical reasons, therefore, it is important to measure fura-2 ratios in GFP-expressing cells with a RFU of <10,000.

**cAMP Level Determination.** For stably transfected lines, cells (4 × 10⁶ cells/well in six-well plates) were incubated overnight in starvation medium (1% serum). After 15-min preincubation with 1 mM 3-isobutyl-1-methylxanthine (IBMX), cells were stimulated with serum-free medium (1% serum). After 15 min of stimulation, media was collected and used to measure the intracellular cAMP levels. The cAMP concentration was determined using a commercial cAMP radioimmunoassay kit to determine cAMP concentrations.

**Confocal Fluorescence Microscopy.** Polylysine-coated 25-mm circular #1 glass coverslips containing live transfected cells were mounted in a modified Leyden chamber containing HEPES-buffered saline solution, pH 7.4. Confocal microscopy was performed on live cells at 22°C using an Olympus Fluoview 2.1 (single-wavelength) or Fluoview 1000 (dual-wavelength colocalization) laser-scanning confocal microscope. Nucleolar localization was marked with Fibulin-HeRed (Fradkov et al., 2002), whereas mitochondrial staining was achieved by prelabeling cells in 25 nM tetramethylrhodamine (TMRM) for 15 min followed by incubation in 5 nM TMRM for the duration of the image collection. Images represent single equatorial planes obtained with a 60× oil objective. Confocal images were processed with Adobe Photoshop 7.0 (Adobe Systems, Mountain View, CA).

**Statistical Analysis.** Unless otherwise stated, data were collected from triplicate wells for each experimental condition. Relative change from baseline data were collected from at least three independent experiments and presented as means ± S.E.M. In calcium signaling experiments, data were collected from n > 30 GFP or RGS2-GFP-expressing individual cells. Statistically significant differences were determined by unpaired Student’s t test method, and a p value of <0.05 was deemed significant. Representative immunoblots shown reflect similar results obtained in at least three separate experiments.

**Results**

**Stable Expression of Human RGS2 mRNA Yielded Multiple Protein Products.** HEK293 cell lines stably transfected with a wild-type RGS2-myc construct expressed multiple RGS2 proteins (30–35-kDa range) compared with empty vector controls (Fig. 1A; compare lanes wt-1 and control). Based on the migration of recombinant RGS2 on SDS-polyacrylamide gel electrophoresis, we predicted that the weak protein band at 34 kDa corresponded to full-length RGS2-myc; however, at least two more highly expressed bands were observed between 30 and 32 kDa. This multiband RGS2 expression pattern was very different from that from cell lines stably transfected with kzRGS2-myc, a construct that was modified to include an optimized translation start consensus [Kozak consensus (Kozak, 1986); kz] RGS2 lines expressed a predominant protein species corresponding to the predicted size of the full-length protein (Fig. 1A; compare lanes kz-1 and kz-2 with wt-1). We examined whether the altered expression pattern correlated with altered signaling function in the wtRGS2 compared with kzRGS2 cell lines. Cell lines with relatively similar expression levels of total RGS2-myc protein (kz-1, 1.0; kz-2, 3.3; and wt-1, 3.7) were used to examine whether the RGS2 expression pattern had an impact on its ability to attenuate Gαq or ACV signaling.

**RGS2-Mediated Inhibition of Gαq Was Similar in kzRGS2- and wtRGS2-Expressing Lines.** To determine whether changes in the RGS2 expression pattern correlated with changes in signaling function, we measured the ability of RGS2 to inhibit Gαq and Gαq/AC signaling in wtRGS2 and kzRGS2 cell lines. We have demonstrated that the Gαq inhibitory function of stably expressed RGS proteins could be compared after stimulation of endogenous muscarinic receptors in HEK293 cells (Heximer et al., 1999; Heximer, 2004). Using a similar assay system, we here compared inositol phosphate accumulation in several RGS2-expressing lines. Basal and carbachol-stimulated inositol phosphate accumulation was lower in all of the RGS2-expressing compared with the control vector-containing cell lines (Fig. 1B) Thus, all of the RGS2 lines studied showed signaling characteristics consistent with the expression of functional RGS2 protein. The level of RGS2-dependent inhibition of inositol phosphate sig-
naling seemed to depend on the levels of total RGS2 protein expression (summed expression of all proteins within 30–35-kDa range) under both basal and carbachol-stimulated conditions. In particular, compared with the control vector cell line, the two high RGS2-expressing lines, wt-1 and kz-2, inhibited the majority of carbachol-stimulated inositol phosphate accumulation, whereas the low RGS2-expressing line kz-1 showed the lowest inhibition of inositol phosphate accumulation (Fig. 1B). Together, these data suggested that RGS2-mediated Goq inhibition is more dependent on the total amount of RGS2 protein in the cell than on differences in its expression pattern.

RGS2-Mediated Inhibition of AC Was Higher in kzRGS2- Compared with wtRGS2-Expressing Lines. HEK293 cells express β-adrenergic receptors that can be stimulated with isoproterenol to increase AC activity and second-messenger intracellular cAMP levels. In Fig. 1C, this pathway was used to determine the relative AC inhibitory activity of RGS2 in wt-1 and kz-2, the two cell lines with the most similar total RGS2 protein levels (Fig. 1A). Isoproterenol stimulation of vector control and wt-1 lines resulted in similar stimulation of cAMP accumulation greater than baseline levels. Data from three independent experiments showed that the wt-1 cell line contained a similarly low level of ACV inhibitory activity as the empty vector control cell line (Fig. 1C). By contrast, the kz-2 cell line showed much less AC-dependent accumulation of cAMP, indicating a higher level of AC inhibitory activity in these cells. Thus, in contrast to the results for Goq inhibition, the inhibition of AC signaling seemed to be highly dependent on the expression of the largest protein species.

Alternative Translation Initiation Produced Four Distinct RGS2 Protein Products. Characterization of the product expression pattern from the endogenous RGS2 gene is difficult because of the lack of antibodies that can reliably detect low levels of protein. Therefore, we constructed an expression reporter construct by cloning the complete 32-base pair 5’-untranslated region and sequences encoding amino acids 1 to 79 of RGS2 in frame ahead of enhanced green fluorescent protein (EGFP) in the pEGFP-C1 vector. The resulting amino-terminal (NT) RGS2-GFP fusion reporter construct, NT-GFP, drove expression of the RGS2 reporter mRNA from the CMV promoter. Transfection of NT-GFP into HEK cells resulted in the expression of four distinct RGS2 protein products compared with nontransfected cells (Fig. 2A, arrows). In agreement with data from stable cell lines (Fig. 1), incorporation of an optimized translation initiation consensus sequence at the first in-frame methionine resulted in production of a single full-length NT-GFP protein (Fig. 2A, lane Kz).

Insertion of an optimized translation initiation consensus sequence at the beginning of the RGS2 open reading frame (ORF) might affect the protein expression pattern by one of two different mechanisms. First, if alternative translation start site use is responsible for the observed multiband profile, then optimization of initiation from the most upstream initiator codon might be expected to reduce the translation start from downstream initiator codons. Second, if the band pattern is due primarily to proteolytic breakdown, a pathway mediated by the position 2 glutamine in RGS2 (Yang et al., 2005; Bodenstein et al., 2007), then mutation of this residue to a stabilizing alanine (required for codon optimization) might stabilize the full-length RGS2 protein and prevent accumulation of smaller breakdown products.

The following observations led us to focus our attention on alternative translation start site use as an explanation for this unique expression profile. Cross-species comparison of human, mouse, and rat RGS2 mRNAs revealed the presence of four conserved in-frame AUG initiator codons that mark the beginning four putative RGS2 ORFs (ORFs 1–4) corresponding to proteins initiated from amino acid positions Met1, Met5, Met16, and Met33 (Fig. 2C). It is noteworthy that the relative migration rates of the four NT-GFP-derived proteins (Fig. 2A) were consistent with translation from four such initiator codons. Alignment of the translation initiation consensus sequence with sequences flanking each putative initiation codons indicated that the third in-frame methionine (Met16) showed the highest degree of sequence similarity to the established translation initiation consensus sequence (Fig. 2B). Moreover, the third-slowest migrating NT-GFP protein was expressed more much strongly than the
others (Fig. 2A), consistent with the possibility of strong relative translation initiation from Met16.

To determine whether the expression profile observed in Fig. 2A was produced by alternative translation initiation, AUG codons at positions Met1, Met5, Met16, and Met33 in NT-GFP were mutated to UUG (leucine) codons, and the resulting protein expression profiles were compared on immunoblots. Ablation of the first two AUG codons, corresponding to Met1 and Met5, completely eliminated expression of full-length and second-most slowly migrating RGS2 bands (Fig. 2D, M1L and M5L). Likewise, ablation of AUG codons at Met16(M16L) and Met33(M33L) selectively abolished expression of the third- and fourth-most slowly migrating protein bands, respectively. Treatment of cells expressing NT-GFP with the proteasome inhibitor MG132 increased expression of all four products but did not dramatically reduce the amount of smaller products, suggesting that the faster-migrating species were not stable byproducts of proteasome-dependent degradation (Fig. 2E). To rule out the unlikely possibility that point mutations in the NT-GFP reporter construct altered the transcription rate or stability of the RGS2 mRNA, reverse transcription-polymerase chain reaction was performed on total RNA samples from cells transfected with the various constructs. Steady-state levels of reporter mRNAs were not different in NT-GFP and NT(M33L)-GFP-transfected cells (data not shown). Thereto
Therefore, data from stable cell lines and mutant NT-GFP translation reporter constructs suggest that the existence of the four protein bands can be explained by alternative translation initiation from four different initiator AUGs corresponding to amino acid positions Met1, Met5, Met16, and Met33 in the RGS2 protein.

**Differential Translation Start Site Model Was Consistent with Stable Cell Line Signaling Data.** Signaling data in Fig. 1 suggest that the RGS2 cDNA can produce a set of proteins that differ in their ability to inhibit AC but not Gαq signaling. The diagram in Fig. 3 summarizes the location of the four putative initiator codons relative to known functional domains in RGS2 (GAP and adenylyl cyclase inhibition, AC, shown below). Figure 3A compares the predicted architecture of proteins expressed in the different stable cell lines from Fig. 1, in which initiation sites are shown as gray shaded bars with forward-facing arrows and are labeled by their amino acid position relative to Met1. Black shaded bars indicate optimization of a translation initiation consensus sequence. Figure 3B compares the wild-type and mutant NT-GFP translation reporter constructs compared in Fig. 2D. We asked whether a single unifying model could explain the relationship between the RGS2 expression pattern and its biological function. Because the kz-1 and kz-2 cell lines apparently express mainly Met1-derived protein compared with Met16-derived protein in the wt-1 line, we inferred that loss of specific sequences between Met1 and Met16 explained the observed lower adenylyl cyclase inhibition by RGS2 in wt-1 cells (Fig. 1). Indeed, the Met16-derived product lacks the AC inhibitory domain (AC) and would be expected to show weaker inhibition of β2-adrenergic signaling than Met1-derived protein (Fig. 3A). Thus, the evidence suggests that alternative translation of RGS2 can produce several RGS2 proteins with different abilities to inhibit adenylyl cyclase activity. Because one test of this model is the functional characterization of each putative RGS2 ORF in isolation, Fig. 3C shows the design of kzORF1 to 4, the expression constructs used for this purpose.

**AC Inhibition Domain Was Not a Key Modulator of RGS2 Plasma Membrane Association.** We showed previously that the RGS2 NTD is required for its association with the plasma membrane and that amphipathic helical sequences between residues 39 and 52 were necessary and sufficient for this function (Heximer et al., 2001). RGS2 can also be found in plasma membrane signaling complexes containing a seven-transmembrane receptor (β2-adrenergic receptor), Gαq, and type IV or VI adenylyl cyclase (Roy et al., 2006a). Thus, it seems that there are multiple discrete domains within the RGS2 NTD that are capable of cooperatively regulating its localization and signaling function. The relative contribution of the AC-inhibition domain to membrane localization is currently unknown and may have important functional implications in cells with different RGS2 expression profiles. Because proteins driven from Met16 and Met33 lack the AC-inhibition domain, the NT-GFP and the AUG-UUG mutant constructs provided a unique opportunity to study the contribution of this domain to membrane association (Fig. 4A). The four NT-GFP products were strongly localized to the plasma membrane with very little GFP fluorescence in the cytoplasm, consistent with the pattern of localization reported previously for the complete amino terminal domain (Heximer et al., 2001). The combined mutation of the first two in-frame AUGs in NT (M1L, M5L)-GFP did not alter tonic plasma membrane targeting efficiency, consistent with the notion that the primary determinants for plasma membrane are located downstream of Met16 in the RGS2 amino terminus (Fig. 4A). Because the AC inhibitory domain is located within amino acids Val9 to His11, it seems unlikely that this domain contributes to basal association of RGS2 with the plasma membrane but rather that it is required for specific recruitment or coordination of activated Gαq-coupled receptor signaling complexes after exposure to a physiologic stimulus.

**Mitochondrial but Not Nuclear/Nucleolar Localization RGS2 Was Dependent on Translation Start Site Use.** It is becoming more widely appreciated that NTD of RGS2 can interact with an increasing number of cellular partners to coordinate localized signaling events (Heximer and Blumer, 2007). Compared with the GFP protein, which is evenly distributed throughout the cytosol and nucleus of HEK293 cells (Heximer et al., 2001), the RGS2 NTD directs nucleoplasmic and possibly nucleolar localization (Fig. 4A). It may be that the cell sequesters RGS2 in the nucleus to prevent its potent inhibition of signaling pathways or that there is a specific purpose for RGS2 inside the nuclear com-

---

**Fig. 3. Schematic representation of cDNA expression constructs showing predicted protein products and domain structures.** Predicted RGS2 protein products expressed from cDNA constructs for myc epitope-tagged proteins used in stable cell lines (A), wild-type and mutant NT-GFP translation initiation reporter (B), and kzORFs (C). All of the predicted protein translation initiation sites are indicated by arrows and amino acid numbers above their corresponding AUG codons shaded in dark gray. Amino acid positions and ORF numbers are indicated relative to the full-length protein sequence (accession number NP_002914). In cases in which initiation codons have been optimized with a Kozak consensus sequence, the shaded AUG codon is highlighted with black. The position of functional domains and protein sequence tags relative to the predicted translation initiation sites are shown below. These are denoted as follows: AC inhibitory domain (AC), plasma membrane-targeting sequence (PM); and GTPase activating protein or RGS core domain (GAP). The 5′ untranslated region of the endogenous RGS2 mRNA has been incorporated into the NT-GFP reporter construct series and is shown above as 5′-UTR. Type and locations of the epitope tags are indicated above the appropriate construct sets and are indicated as follows: GFP and triple myc epitope tag (3×myc).
partment. We therefore examined its localization to identify new potential sites of RGS2 function. The NT-GFP proteins showed strong colocalization (arrows) with the nucleolar marker Fibrillarin-HcRed (Fig. 4B), indicating a possible role for RGS2 in nucleoli. NT-GFP-derived proteins also associated with punctate organelar structures in the cytosol predicted previously to be mitochondria (Heximer et al., 2001). These features were shown to precisely colocalize (arrowheads) with the mitochondrial-specific dye TMRM (Fig. 4C). It is the only Met33-initiated protein that targets mitochondria, because TMRM colocalization was abolished for the NT(M33L)-GFP construct. It remains to be determined whether RGS2, and more specifically its Met33-derived ORF, plays a role in the regulation of mitochondrial function. kzORF1 to 4 are recruited from the nucleus by Goq, but not Goq/ACV signaling. Although the RGS2 NTD mediates localization and AC inhibition, the RGS2 GAP domain mediates its function as a Goq inhibitor. Our group and others have shown that these protein domains cooperate to mediate recruitment of RGS2 from the nucleus in response to a Goq stimulus (Heximer et al., 2001; Roy et al., 2003). We asked whether long-term Goq/ACV signaling can also recruit these four RGS2 proteins (kzORFs) out of the nucleus (Fig. 5). We predicted that RGS2 proteins containing the AC-inhibition domain (kzORF1 and kzORF2) would be more efficiently recruited to the plasma membrane. Expression of each RGS2 product was achieved by polymerase chain reaction cloning and inclusion of an optimized translation initiation sequence at the upstream AUG codon (Fig. 3C) The resulting clones were named kzORF1 through kzORF4. Each kzORF construct expressed a predominant protein band on anti-GFP immunoblots (Fig. 5A). Confocal microscopy was used to examine the subcellular localization of kzORF1 to -4 in control cells and in cells coexpressing constitutively active Goq or Goq/ACV (Fig. 5, B and C). All four kzORF clones showed efficient recruitment from the nucleus to the plasma membrane/cytosol compartment in response to Goq activation (Fig. 5B). By contrast, none of the different kzORF constructs tested was efficiently recruited from the nucleus to the plasma membrane/cytosol in Goq/ACV-stimulated cells. Relative pixel intensity values indicated that Goq activation resulted in recruitment of kzORF1 from the nucleus, whereas Goq/ACV activation had no effect on its relative distribution, despite the presence of an intact AC inhibition domain (Fig. 5C).

Taken together, the subcellular localization data for the NT-GFP and kzORF constructs do not support a role for

![Fig. 4. Subcellular localization of RGS2 amino terminal domains produced from wild-type and mutant translation initiation reporter constructs. A, localization of wild-type and indicated NT-GFP mutation constructs were analyzed in transfected living HEK cells using confocal microscopy. Images show cells with low/medium relative fluorescence and are representative of at least 50 cells transfected with the same construct. Confocal images were taken of HEK cells transfected with NT-GFP. Shown are GFP images collected from the basal region of the cell as determined by a z-axis series. B, colocalization of NT-GFP constructs with nucleolar markers in live cells. HEK cells were cotransfected with wild-type NT-GFP and the nucleolar marker protein fibrillarin (Fibrillarin-HcRed). Using different lasers for excitation (488 nm, GFP; 543 nm, HcRed) and emission spectrum discrimination capabilities of the Olympus Fluoview confocal microscope, green- and red-channel images were collected from the same confocal plane to determine the subcellular localization of NT-GFP and Fibrillarin-HcRed, respectively. Merged images were created to demonstrate the extent of colocalization (yellow) of these constructs. C, colocalization of NT-GFP constructs with mitochondrial dyes in live cells. HEK cells transfected with the wild-type or M33L NT-GFP constructs were incubated in the mitochondrial targeted fluorescent dye TMRM. Colocalization was determined as in B.](image)

![Fig. 5. Effect of GAP domain function and G-protein-signaling status on RGS2 localization determinants. A, Western blot of total cell lysate from cells transfected with the specified construct shows that the presence of a Kozak consensus sequence results in the production of only one protein species. B, localization of the indicated RGS2 kzORF-GFP fusion constructs with and without either constitutively active Goq(Gq*) or Goq and ACV(Gq*/ACV) was examined in transiently transfected cells as described above. C, the ratio of GFP signal between the nucleus and plasma membrane was measured using ImageJ. Shown are means ± S.E.M.](image)
alternative translation initiation in the differential control of RGS2 targeting to the plasma membrane or recruitment from the nucleus. Therefore, functional differences between the RGS2 ORFs are most likely to be the result of their intrinsic inability to inhibit Goq or ACV.

KzORFs 1 to 4 Showed Similar Levels of Goq Inhibitory Function. Data from our translation reporter system and RGS2 stable lines suggest that all four RGS2 proteins produced by alternative translation were functionally competent with respect to their Goq inhibition activity. However, it was not possible to determine the relative function of the individual products because these proteins were expressed simultaneously from the wild-type RGS2 mRNA construct. First, we examined Goq inhibition by each individual RGS2 ORF under long- and short-term signaling conditions. We measured the ability of kzORF1 to -4 to inhibit inositol phosphate accumulation in cells cotransfected with constitutively active Goq(R183C). Transfection of Goq(R183C) alone resulted in a ~50-fold increase in accumulated inositol phosphate (IPx) levels relative to nontransfected HEK293 cells. It is noteworthy that Goq(R183C)-dependent phosphoinositide hydrolysis was attenuated to a similar extent (>80% reduction of maximum signal) by each of the different RGS2 proteins (Figs. 6A). In a separate series of experiments in which 3-fold less RGS2 plasmid was used, kzORF1 and kzORF3 both attenuated signaling to a similar extent (~40% reduction of maximal signal; data not shown). In short-term assays, HEK293 cells stably expressing the M1 muscarinic receptor (M1-HEK) were used to study the function of the various RGS2 ORFs 1 to 4 as inhibitors of agonist-mediated increases in intracellular calcium. In particular, M1-HEK cells were transiently transfected with pEGFP control plasmid or the indicated RGS2 kzORF-GFP fusion construct before fura-2 loading and stimulation with carbachol. The Goq inhibitory function of ORFs 1 to 4 was determined by measuring intracellular calcium responses in single cells that had been preselected on the basis of their kORF-GFP expression. When RGS2 activity was compared between cells expressing similar levels, the four kzORFs all showed similar inhibition of intracellular calcium elevation (~40%) in response to a 200 μM carbachol bolus (Fig. 6B). Taken together, these data indicated that the four RGS2 ORFs produced by alternative translation initiation were not functionally different at the level of their Goq inhibition.

Alternative RGS2 Translation Start Sites Produced Functionally Distinct Inhibitors of AC. Data from Fig. 1C suggest that the different RGS2 proteins produced by alternative translation of the RGS2 mRNA may behave differently in their abilities to attenuate G-protein-coupled receptor-mediated cAMP accumulation in HEK cells. We predicted that these differences were attributed to the specific loss of the AC inhibitory domain in Met16 (ORF3)- and Met33 (ORF4)-derived proteins. To determine the relative AC inhibitory potential of the various alternatively translated proteins, we used a cotransfection assay that was developed to study the function of the RGS2 as a direct inhibitor of ACV function (Salim et al., 2003). The various kzORFs were transiently cotransfected with constitutively active Gs(Q227L) and ACV in HEK293 cells, after which cAMP accumulation was measured (Fig. 7). In the presence of active Gs(Q227L) and ACV, steady-state intracellular cAMP levels were increased by ~20-fold compared with unstimulated controls. The coexpression of the full-length RGS2 (kzORF1) and kzORF2 proteins each resulted in a >50% decrease in cAMP levels. As predicted from the expected downstream initiator codon positions relative to the AC inhibitory domain, kzORF3 and kzORF4 were completely deficient at inhibiting cAMP accumulation by Gs(Q227L) and ACV.

Activation of Goq but Not Goq Signaling Pathways Altered the Expression Profile of RGS2 Translation Products. Because the above data show that the biological activity of RGS2 depends on the relative expression levels of different proteins produced from different translation initiation sites, we next determined whether the relative abundance of the alternative translation products was regulated by different long-term G-protein signaling conditions (Fig. 8). Therefore, the translation reporter vector NT-GFP was expressed alone or together with either Goq(R183C) or Goq(Q227L) and ACV. The resulting protein expression profiles were compared on immunoblots. Although no changes in the RGS2 profile were observed in response to Goq stimulation, coexpression of Goq and ACV resulted in an increase in the relative abundance of kzORFs 1 to 4 as inhibitors of agonist-mediated increases in intracellular calcium (Fig. 6B). Taken together, these data indicated that the four RGS2 ORFs produced by alternative translation initiation were not functionally different at the level of their Goq inhibition.
the expression of the Met1-derived protein (Fig. 8A). Also evident was a concurrent decrease in the expression of the Met16-derived protein such that the ratio of Met1 to Met16-derived protein was greatly increased in response to Goα and ACV (Fig. 8B). Because the transcriptional activity of both the RGS2 and CMV promoters is increased in a cAMP-dependent manner, we asked whether the observed increase in the Met1-derived protein was caused by transcriptional up-regulation after cotransfection with Goα and ACV. The NT-GFP cassette was cloned into a tetracycline-inducible vector (pREV-TRE), and the RGS2 expression profile was examined at different rates of transcription that were controlled by the amount of doxycycline added to the culture medium. Increased transcription from this reporter construct was evident from stepwise increases in RGS2 protein expression; however, this was not associated with an increased level of the Met1-derived compared with the Met16-derived protein (Supplemental Data). Moreover, the coexpression of Goα and ACV resulted in increased relative expression of the Met1-derived protein irrespective of the doxycycline concentration used. Together, these data suggest that the Met1-derived protein is up-regulated independently from cAMP-dependent changes in transcription rate.

**Discussion**

**RGS Protein Genes Expressed Multiple Gene Products with Different Functional Properties.** As completion of the human and mouse genome sequencing projects draw near, the search for novel RGS protein products with different biological functions is an emerging area of interest. Alternative mRNA splicing is a common mechanism by which several RGS proteins are produced from a single gene. Genes such as RGS3 and RGS12 produce alternatively spliced mRNAs that furnish their respective GAP domains with varying complements of PDZ domain or PDZ domain binding sequences (Snow et al., 1998; Kehrl et al., 2002). Likewise, RGS6, RGS8, RGS9, RGS10, and RGS11 yield spliced variants of their GAP domain sequences with more than one complement of regulatory domains (Graneman et al., 1998; Giudice et al., 2001; Haller et al., 2002; Saitoh et al., 2002; Chatterjee et al., 2003). It is intriguing that nearly all of the RGS proteins derived from alternatively spliced mRNAs contain the RGS GAP domain sequences. Thus, it seems that cells modulate their G-protein signaling profiles via alternative splicing of appropriate regulatory domains onto RGS domain sequences. It is of interest, therefore, to characterize the mechanisms for alternative RGS protein production as a step toward understanding cellular modulation of G-protein signaling.

**Multiple RGS2 Proteins Were Expressed from a Single mRNA.** Our analysis showed that RGS2, like many of the RGS protein-encoding genes, is capable of producing more than one protein product. By contrast, however, RGS2 did not seem to use differential splicing to generate these species. Some RGS proteins (RGS2, RGS4, RGS5, and RGS16) are targeted for proteasome-mediated degradation through the N-end rule pathway, a mechanism that is dependent on cleavage of the first methionine and the protein stabilizing/destabilizing nature of the second amino acid (Davydov and Varshavsky, 2000; Hu et al., 2005; Lee et al., 2005; Bodenstein et al., 2007). We asked whether such a mechanism could produce the observed expression profile. Our current data do not seem to support this notion. First, according to the eukaryotic N-end rule (Varshavsky, A., 1996), the glutamine, phenylalanine, aspartic acid, and lysine residues at position 2 (Table 1) are all destabilizing residues and therefore should not promote selective accumulation of any of the four species. Second, the addition of the proteasome inhibitor MG132 to cells expressing NT-GFP did not selectively stabilize the full-length RGS2 band at the expense of the smaller proteins. Although we cannot rule out the possibility that some of the ORF1- to -4-derived proteins have a
higher intrinsic stability than the others, the specific loss of single protein bands after AUG mutagenesis clearly points to alternative translation initiation as the primary explanation for the four species.

**Alternative Translation Initiation Yields RGS2 Proteins with Varying Biological Functions.** To the best of our knowledge, this is the first example of alternative translation initiation leading to the expression of RGS protein products capable of conferring different biological activities. Indeed, the abilities of the different RGS2 products to inhibit AC were dramatically different. Consistent with their lack of an AC inhibitory domain, kzORF3 and kzORF4 were deficient of AC inhibitory function. These data provide supporting evidence for the notion that alternative translation start site initiation is another potential mechanism for the regulation of RGS protein function in mammalian cells. It is of interest to determine whether these alternatively translated proteins are capable of differentially regulating other recently identified RGS2 effectors such as TRPV6, for which the interaction domain in RGS2 amino terminus is not known.

**Leaky Ribosome Scanning Promotes Alternative Translation Initiation of RGS2.** Our data support the use of four different initiation codons in the RGS2 mRNA and predict that a number of ribosomes are able to bypass the upstream initiator codon(s) in the wild-type RGS2 mRNA. Three mechanisms have been proposed to explain how translation from multiple ORFs in a single mRNA is achieved: 1) internal ribosome entry; 2) ribosome shunting; and 3) leaky ribosome scanning (Kozak, 1991). Internal ribosome entry has been described for a number of genes, including c-myc and the p58 and p110 PITSLRE protein kinases (Nanbru et al., 1997; Cornelis et al., 2000). Inclusion of a strong translation initiation signal at the first in-frame AUG codon resulted in loss of expression of the other smaller RGS2 products. Although this mutant also incorporates a stabilizing alanine residue at the second amino acid position that could increase its relative stability compared with the other ORFs, this mechanism cannot explain the loss of expression of the other products. Moreover, the loss of specific protein bands in the M>L mutagenesis experiments suggests that translation start site use is the primary factor controlling the observed change in expression pattern.

What is the mechanism controlling differential translation start site use in the RGS2 cDNA? Ablation of downstream ORF initiation in the RGS2-myc cell lines and the cells expressing the NT-GFP reporter argues strongly against the possibility that the RGS2 mRNA contains one or more strong internal ribosome entry site elements. At present we cannot rule ribosome shunting on the RGS2 mRNA, a mechanism in which ribosomes are repositioned across strong RNA hairpins. However, this mechanism requires termination of translation of the upstream ORF and reinitiation of the shunted ribosomes (Hemmings-Miesczak et al., 2000). The multiple RGS2 ORFs in question are long overlapping sequences that would make a termination/reinitiation event via ribosome shunting highly improbable.

In the ribosome scanning model of translation, the 43S ribosomal complex scans the 5'-UTR in a 5' to 3' direction until it reaches an AUG within the context of a good consensus initiation sequence, where translation begins. Leaky ribosome scanning can produce multiple protein products if the translation machinery does not efficiently recognize the upstream initiator codons. To date, only a small subset of cellular mRNAs has been reported to express protein products from more than one start codon. Among these are CCAAT/enhancer-binding proteins α and β that each give rise to multiple products because of weak translation initiation consensus sequences at their upstream AUG codons (Calkhoven et al., 2000). One predicted consequence of the leaky scanning model is that initiation from downstream AUG codons should increase if upstream alternative start sites are disrupted. Indeed, our studies with the NT-GFP translational reporter show that this occurs in the RGS2 mRNA because disruption of the initiator sequence for the strongly recognized AUG codon for Met16 results in increased expression of the Met33-derived protein. Thus, it seems likely that RGS2 can be added to this small set of genes whose protein expression profile is mediated by leaky ribosome scanning.

**G-Protein Signaling Status Regulates the RGS2 Protein Expression Profile.** RGS2 gene expression is highly tuned to the signaling status of the cell. RGS2 is an immediate-early gene whose mRNA levels are increased in a number of cell types in response to stimuli that increase intracellular calcium and cAMP (Kehrl and Sinnarajah, 2002). Thus, it has been proposed that RGS2 mRNA levels may be increased as part of a negative feedback mechanism to reciprocally modulate Gαq- and AC-dependent signaling (Roy et al., 2006b). The current study suggests that the regulation of post-transcriptional events may be equally important for integrating signaling feedback loops. In particular, long-term Gαq signaling increased expression of the Met1-derived compared with the Met16-derived protein, an observation consistent with signaling-dependent modulation of translation efficiency at Met1. It is noteworthy that in forskolin-treated osteoblasts, the largest RGS2 protein is also apparently expressed at much higher levels than the other proteins, suggesting that a similar regulatory mechanism may be present in other cell types and tissues (Roy et al., 2006b). The precise mechanism for this unique type of regulation, however, remains to be determined. Nonetheless, this unique adaptation of the RGS2 expression profile to a change of cell signaling status represents a new type of signaling feedback mechanism that implicates regulated alternative translation start site use in the regulation of G-protein-coupled signaling.

### Table 1

<table>
<thead>
<tr>
<th>Amino terminal sequence of ORFs 1 to 4 derived from the human RGS2 gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>The first 14 amino acids of each predicted RGS2 ORF are shown. Amino acid positions correspond to the full-length protein sequence (accession number NP_002914). Second-position residues glutamine, leucine, aspartic acid, and lysine of each ORF are highlighted in boldface type.</td>
</tr>
</tbody>
</table>

| 1M | Q | S | A | M | F | L | A | V | Q | H | D | C | R | 14 | ORF1 |
|----|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|------|
| 4M | L | A | V | Q | H | D | C | R | P | M | M | D | K | 17 | ORF2 |
| 16M | D | K | S | A | G | S | G | H | K | S | E | E | K | 20 | ORF3 |
| 33M | K | R | T | L | L | K | D | W | K | T | R | L | S | 46 | ORF4 |