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DAPLE protein inhibits nucleotide exchange on $G\alpha_s$ and $G\alpha_q$ via the same motif that activates $G\alpha_s$

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Besides being regulated by G-protein-coupled receptors, the activity of heterotrimeric G proteins is modulated by many cytoplasmic proteins. GIV/Girdin and DAPLE (Dvl-associating protein with a high frequency of leucine) are the best-characterized members of a group of cytoplasmic regulators that contain a $G\alpha$ -binding and -activating (GBA) motif and whose dysregulation underlies human diseases, including cancer and birth defects. GBA motifcontaining proteins were originally reported to modulate G proteins by binding $G\alpha$ subunits of the $G_{i/o}$ family $(G\alpha_i)$ over other families (such as G_s , $G_{q/11}$, or $G_{12/13}$), and promoting nucleotide exchange in vitro. However, some evidence suggests that this is not always the case, as phosphorylation of the GBA motif of GIV promotes its binding to $G\alpha_s$ and inhibits nucleotide exchange. The G-protein specificity of DAPLE and how it might affect nucleotide exchange on G proteins besides $G\alpha_i$ remain to be investigated. Here, we show that DAPLE's GBA motif, in addition to $G\alpha_i$, binds efficiently to members of the G_s and $G_{\alpha/11}$ families ($G\alpha_s$ and $G\alpha_{\alpha'}$) respectively), but not of the $G_{12/13}$ family $(G\alpha_{12})$ in the absence of post-translational phosphorylation. We pinpointed Met-1669 as the residue in the GBA motif of DAPLE that diverges from that in GIV and enables better binding to $G\alpha_s$ and $G\alpha_g$. Unlike the nucleotide-exchange acceleration observed for $G\alpha_{i}$, DAPLE inhibited nucleotide exchange on $G\alpha_s$ and $G\alpha_g$. These findings indicate that GBA motifs have versatility in their G-protein-modulating effect, i.e. they can bind to $G\alpha$ subunits of different classes and either stimulate or inhibit nucleotide exchange depending on the G-protein subtype.

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This article contains Figs. S1-S4.

Heterotrimeric G proteins are essential signaling molecules that relay extracellular signals acting on G-protein-coupled receptors (GPCRs)⁴ to intracellular effector proteins (1). They are involved in a plethora of physiological processes and mediate the effect of >30% of Food and Drug Administrationapproved drugs (2). At the molecular level, heterotrimeric G proteins switch between "on" or "off" states depending on their guanine nucleotide-binding status. In the inactive state, GDPbound $G\alpha$ subunits associate tightly with obligate $G\beta\gamma$ dimers, which in turn serve to prevent spurious nucleotide exchange by working as guanine nucleotide-dissociation inhibitors (GDIs) (3, 4). The $G\alpha\beta\gamma$ trimer is the substrate for the guanine nucleotide-exchange factor (GEF) activity of GPCRs, which promote the exchange of GDP for GTP on $G\alpha$ and the subsequent dissociation of $G\beta\gamma$. Both $G\alpha$ -GTP and free $G\beta\gamma$ become active signaling species that engage their own set of effectors to propagate downstream signaling. $G\alpha$ subunits are classified into four families depending on sequence conservation and the effector targets that they modulate: G_s , $G_{i/o}$, $G_{q/11}$, and $G_{12/13}$ (5, 6). Signaling is terminated upon GTP hydrolysis by $G\alpha$, which leads to reassociation with $G\beta\gamma$ to form again an inactive $G\alpha\beta\gamma$ trimer.

Based on the above, it is evident that the amplitude and duration of G-protein signaling is highly dependent on the regulation of nucleotide handling by $G\alpha$. In this regard, it has become increasingly clear that G-protein activity is controlled by a complex network of regulators that expands beyond GPCRs and $G\beta\gamma$. The best-characterized ones are the regulators of G-protein signaling (RGS) proteins, which are GTPase-accelerating proteins (7–16). RGS proteins, as well as some effectors like PLC- β isoforms, enhance the rate of nucleotide hydrolysis on $G\alpha$, thereby facilitating the termination of G-protein signaling (16, 17). Another group of regulators, defined by the presence of a sequence called the GOLoco motif, locks $G\alpha$ in the GDP-

⁴ The abbreviations used are: GPCR, G-protein–coupled receptor; GBA, $G\alpha$ -binding and -activating; GEF, guanine–nucleotide exchange factor; GDI, guanine nucleotide–dissociation inhibitor; RGS, regulator of G-protein signaling; GIV, $G\alpha$ -interacting, vesicle-associated protein; DAPLE, Dvl-associating protein with a high frequency of leucine; PLC, phospholipase C; GST, glutathione S-transferase; IP, immunoprecipitation; IB, immunoblotting; GTPγS, guanosine 5'-O-(thiotriphosphate); DMEM, Dulbecco's modified Eagle's medium; IPTG, isopropyl β -D-1-thio-galactopyranoside; PMSF, phenylmethylsulfonyl fluoride; PVDF, polyvinylidene difluoride; PDB, Protein Data Bank; PBM, PDZ-binding motif; aa, amino acid.



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bound state by virtue of their GDI activity (18-26). There are also nonreceptor proteins that have the same biochemical activity as GPCRs, i.e. they are GEFs (27-32). Among them, it has been possible to link the GEF activity to a defined protein domain or sequence only for a subset of these nonreceptor G-protein regulators. This is for proteins that contain a $G\alpha$ -binding and -activating (GBA) motif, an evolutionarilyconserved sequence of ~30 amino acids with a well-defined mechanism of action at the structural level (32-37). Six GBA motif-containing proteins have been identified to date: GIV, DAPLE, Calnuc, NUCB2, PLCδ4b, and the C. elegans protein GBAS-1 (32, 33, 38–40).

Among the GBA motif-containing proteins, GIV (also known as Girdin) and DAPLE were the first ones to be identified and are the best-characterized ones to date (32, 34, 35, 39, 41). Early evidence indicated that both proteins bind to inactive, GDP-bound $G\alpha_i$ subunits $(G\alpha_{i1}, G\alpha_{i2}, \text{ and } G\alpha_{i3})$ to accelerate their rate of spontaneous nucleotide exchange in vitro, although they interacted poorly with other $G\alpha$ subunits, including other members of the $G_{i/o}$ family like $G\alpha_o$ (39, 42). This biochemical activity in vitro correlates well with an enhancement of G-protein signaling observed in cells. For example, it has been shown that the GBA motif of these proteins is required for signaling readouts that depend on $G\alpha_i$ (like inhibition of adenylyl cyclase or antibodies that specifically detect GTP-bound $G\alpha_i$) (39, 40, 43–45), as well as on $G\beta\gamma$ (like free G $\beta\gamma$ biosensors or the PI3K–Akt effector pathway) (32, 39-41, 44, 46-48). It has also been recently shown that G-protein regulation by the GBA motif of DAPLE operates in vivo to control vertebrate development (41, 49). Overall, the biomedical relevance of this signaling mechanism is highlighted by multiple studies establishing the involvement of G-protein regulation by GIV and DAPLE in several human disorders like cancer, liver fibrosis, or embryonic defects (39, 41, 45–47, 50).

Although the seminal studies described above indicated that the GBA motif—containing proteins are GEFs that specifically work on $G\alpha_i$ subunits, more recent evidence has challenged this notion by suggesting that the biochemical activity and G-protein specificity of GBA motifs can vary from this. More specifically, it has been shown that upon sequential phosphorylation of the GBA motif of GIV at serine 1674 and serine 1689 by two different kinases, it does not bind to $G\alpha_i$ proteins anymore (51). Instead, such phosphoevents enhance the affinity of GIV's GBA motif for $G\alpha_s$, which is poor in the absence of phosphorylation (51, 52). Interestingly, this interaction results in the inhibition of nucleotide exchange rather than in acceleration as observed for $G\alpha_i$ (51). Together, these findings suggest that GIV can *switch* from behaving as a GEF (for $G\alpha_i$) to behaving as a GDI (for $G\alpha_s$) upon phosphorylation. The term guanine nucleotide– exchange modulator has been proposed for GIV to convey that it can have different effects on nucleotide handling depending on context (51). However, it is not known whether any GBA motif present in a protein can have the simultaneous ability to work as a GEF or GDI or to interact efficiently with G α subunits besides $G\alpha_i$, in the absence of phosphomodifications. Here, we set out to characterize the previously unexplored G-protein selectivity of DAPLE, and we found that its GBA motif can efficiently bind to representative members of three different G-protein subfamilies $(G_{i/o}, G_s, \text{ and } G_{q/11})$ without any phosphomodification. For $G\alpha_s$ and $G\alpha_g$, this interaction results in the inhibition of nucleotide exchange, suggesting that DAPLE can at the same time work as a GEF or as a GDI depending on the G-protein substrate.

Results

DAPLE binds efficiently $G\alpha_s$ and $G\alpha_a$ in addition to $G\alpha_{i3}$

We set out to characterize the G-protein specificity of DAPLE by investigating its binding to representative $G\alpha$ subunits of each one of the four different families, i.e. $G\alpha_{i3}$ from $G_{i/o}$, $G\alpha_s$ from G_s , $G\alpha_q$ from $G_{q/11}$, and $G\alpha_{12}$ from $G_{12/13}$. For this, we carried out protein-protein-binding experiments using lysates of HEK293T cells expressing each one of the G proteins and GST-fused DAPLE immobilized on resin. As an internal reference for these experiments, we used GST-GIV, which has been previously shown to have a marked preference for $G\alpha_i$ over other G proteins. The two GST-fused constructs consisted of C-terminal fragments of each protein containing the GBA motif (GST-DAPLE aa 1650-2028 and GST-GIV aa 1671–1755, see Fig. 1A). Purification of any of the two constructs based on affinity capture of their N-terminal GST tags resulted in proteins with degradation products that could not be avoided despite attempts to optimize expression conditions. However, all or most of the degradation products should contain G-protein-binding sites because the GBA motif is adjacent to the GST tag used for affinity capture (Fig. 1A). Based on this, we reasoned that using equal amounts of total GST-fused proteins would allow the direct comparison of G-protein binding to GST-DAPLE and GST-GIV because the number of binding sites should be similar. If so, the prediction was that we should see equivalent binding of $G\alpha_{i3}$ to GST-DAPLE and GST-GIV because it has been previously reported that these two constructs have equivalent affinity for this G protein (39). We found that this is the case because the same amount of $G\alpha_{i3}$ FLAG was detected in resin-bound complexes of GST-DAPLE and GST-GIV (Fig. 1B, left panel).

Having established these experimental conditions to semiquantitatively assess G-protein binding to DAPLE compared with GIV, we performed equivalent experiments with $G\alpha_s$, $G\alpha_a$, and $G\alpha_{12}$. As expected, binding of $G\alpha_s$, $G\alpha_a$, or $G\alpha_{12}$ to GST-GIV was undetectable or marginal relative to the binding observed for $G\alpha_{i3}$ (the same proportion of input lysate was run in each experiment to facilitate the comparison of relative binding across G proteins) (Fig. 1B). In contrast, we detected robust binding of $G\alpha_s$ and $G\alpha_g$, but not $G\alpha_{12}$, to GST–DAPLE, which was comparable with the binding observed for $G\alpha_{i3}$ based on comparison with their respective input lanes. To rule out that the observed binding of $G\alpha_s$ and $G\alpha_g$ to GST–DAPLE was due to the overexpression of G proteins, we performed analogous experiments to detect binding of $G\alpha_s$ or $G\alpha_q$ endogenously expressed in HEK293T cells. We found that endogenous $G\alpha_{\rm s}$ and $G\alpha_{a}$ bind efficiently to GST-DAPLE but not to GST-GIV (Fig. S1). These results indicate that, in contrast to GIV, nonphosphorylated DAPLE can bind robustly to G proteins from three different subfamilies: $G_{i/o}$ ($G\alpha_{i3}$), G_s ($G\alpha_s$), and $G_{\alpha/11}$ ($G\alpha_a$).



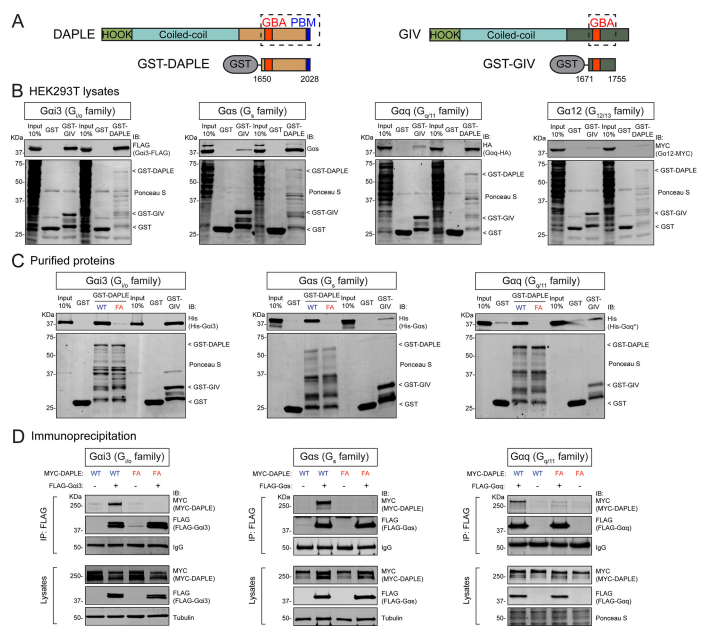


Figure 1. DAPLE binds efficiently to $Gα_s$ and $Gα_q$ through its GBA motif. *A*, bar diagrams depicting the domains of DAPLE (left) and GIV (right) and the fragments of each one fused to GST used for experiments shown in this figure. *B*, DAPLE binds efficiently to $Gα_{i3}$, $Gα_s$, and $Gα_q$ but not to $Gα_{12}$, whereas GIV only binds efficiently to $Gα_{i3}$ among the G proteins tested. Lysates of HEK293T cells transfected with $Gα_{i3}$ –FLAG, $Gα_s$, $Gα_q$ –HA, and $Gα_{12}$ –MYC were incubated with GST, GST–DAPLE, or GST–GIV immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. *C*, DAPLE WT but not DAPLE F1675A (*FA*) binds to purified $Gα_{i3}$, $Gα_s$, and $Gα_q$. His– $Gα_{i3}$, His– $Gα_s$, or His– $Gα_q^*$ were incubated with GST, GST–DAPLE (WT or FA mutant), or GST–GIV immobilized on GSH-agarose beads, and bead-bound proteins were detected by Ponceau S staining or IB as indicated. *D*, DAPLE WT, but not DAPLE FA mutant, co-immunoprecipitates with $Gα_{i3}$, $Gα_s$, and $Gα_q$. Lysates of HEK293T cells co-expressing full-length MYC–DAPLE (WT or FA mutant) with the indicated FLAG-tagged G proteins (or no tagged G protein as negative control) were subjected to IP with a FLAG antibody, and bound proteins were detected by IB as indicated. The lower immunoblot panels (lysates) correspond to aliquots of the starting material used for IPs shown in the upper panels (IP: FLAG). All results presented in this figure are representative of at least three independent experiments ($n \ge 3$).

DAPLE binds directly to $G\alpha_s$ and $G\alpha_q$ via its GBA motif

Next, we investigated whether the association of DAPLE with $G\alpha_s$ and $G\alpha_q$ described above is mediated by direct binding of the G proteins to the GBA motif of DAPLE as observed previously for $G\alpha_i$ (39). For this, we carried out experiments with purified G proteins instead of cell lysates, and we included a GST–DAPLE construct bearing a mutation in its GBA motif (F1675A) that has been previously shown to disrupt binding to $G\alpha_{i3}$ (39). We also kept GST–GIV as an internal control in these experiments. The G proteins were His-tagged versions of

 $G\alpha_{i3}$, $G\alpha_{s}$, and $G\alpha_{q}$ purified from bacteria. For $G\alpha_{q}$, we used a chimera containing partial sequences of $G\alpha_{i}$ (named here $G\alpha_{q}^{*}$) that can be expressed in bacteria and that have been previously validated to bind to a wide range of $G\alpha_{q}$ -specific partners (53). As observed with G proteins from cell lysates, both GST-DAPLE and GST-GIV bound similarly to $G\alpha_{i3}$, but only GST-DAPLE bound efficiently to $G\alpha_{s}$ and $G\alpha_{q}^{*}$ (Fig. 1C). DAPLE binding to any of the G proteins was disrupted by the F1675A mutation (Fig. 1C), indicating that the GBA motif of DAPLE is required for the direct binding of $G\alpha_{s}$ and $G\alpha_{q}$.

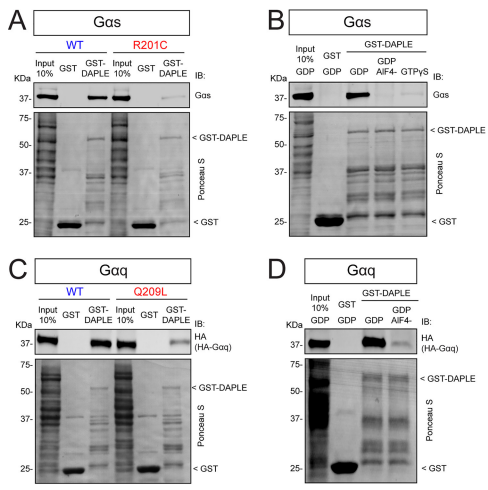


Figure 2. DAPLE binds preferentially to inactive versus active $G\alpha_s$ or $G\alpha_q$. A, binding of DAPLE to the constitutively-active $G\alpha_s$ mutant R201C is diminished compared with $G\alpha_s$ WT. Lysates of HEK293T cells expressing $G\alpha_s$ WT, or $G\alpha_s$ R201C were incubated with GST or GST–DAPLE immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. B, binding of DAPLE to $G\alpha_{\varsigma}$ loaded with $GDP \cdot AlF_{4}^{-}$ or with $GTP \cdot YS$ is diminished compared with binding to $G\alpha_s$ loaded with GDP. Lysates of HEK293T cells expressing $G\alpha_s$ were incubated with nucleotides as indicated under "Experimental procedures" and incubated with GST or GST-DAPLE immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. C, binding of DAPLE to the constitutively-active $G\alpha_q$ mutant Q209L is diminished compared with $G\alpha_q$ WT. Lysates of HEK293T cells expressing $G\alpha_q$ -HA WT or $G\alpha_q$ -HA Q209L were incubated with GST or GST-DAPLE immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. D, binding of DAPLE to $G\alpha_q$ loaded with GDP_4^- is diminished compared with binding to $G\alpha_q$ loaded with GDP_4^- Lysates of HEK293T cells expressing $G\alpha_q$ were incubated with nucleotides as indicated under Experimental procedures" and incubated with GST or GST-DAPLE immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. All results presented in this figure are representative of two independent experiments (n = 2).

To determine whether the binding of $G\alpha_s$ and $G\alpha_g$ to a fragment of DAPLE in vitro observed above occurs when the GBA motif is in the context of the full-length protein expressed in cells, we carried out co-immunoprecipitation experiments (co-IP) from HEK293T cells. FLAG-tagged $G\alpha_{i3}$, $G\alpha_{s}$, or $G\alpha_{q}$ was co-expressed with MYC-tagged full-length DAPLE, and IPs were carried out with FLAG antibodies. Cells expressing MYC-DAPLE in the absence of FLAG-G α were used as negative control. We found that MYC-DAPLE WT was efficiently co-IPed by FLAG-G α_{i3} , FLAG-G α_{s} , or FLAG-G α_{q} (Fig. 1D). Moreover, we found that the GBA motif mutant F1675A prevents the co-IP of DAPLE with any of the three G proteins (Fig. 1D). These findings indicate that full-length DAPLE interacts with $G\alpha_s$ and $G\alpha_q$ through its GBA motif, much like $G\alpha_i$ proteins do.

DAPLE binds to inactive but not active $G\alpha_s$ and $G\alpha_a$

A property of all previously characterized GBA motifs is that they bind preferentially to inactive (GDP-bound) but not to active (GTP-bound) conformations of $G\alpha_i$ (32, 33, 37–40). Thus, we set out to test whether this is also the case for the DAPLE- $G\alpha_s$ and DAPLE- $G\alpha_g$ interactions. For this, we first compared DAPLE binding to constitutively active, GTPase-deficient mutants of $G\alpha_s$ and $G\alpha_g$ (R201C and Q209L, respectively (54)) versus their WT G-protein counterparts. We found that both active mutants have diminished binding to DAPLE compared with WT (Fig. 2, A and C). As a complementary approach to address this point, we performed similar experiments with $G\alpha$ subunits loaded with nucleotides that mimic different G-protein activation states. More specifically, G proteins were loaded with GDP (inactive conformation), GDP·AlF₄ (which mimics the GTP-bound activation transition state) or GTP γ S (a nonhydrolyzable GTP analog). The GTP γ S condition was excluded for $G\alpha_q$ because it is known that this G protein exchanges nucleotide very slowly and loads GTPγS substoichiometrically even under experimental conditions that

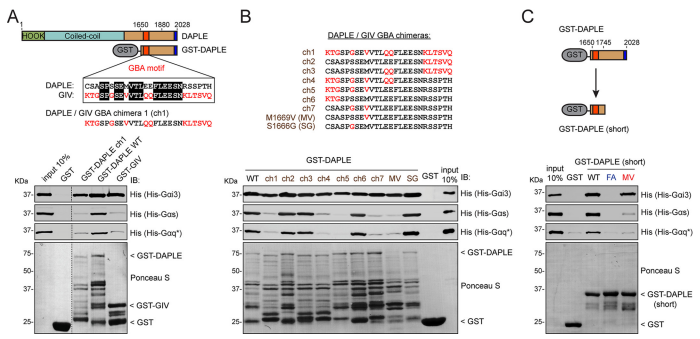


Figure 3. Met-1669 in DAPLE is responsible for its enhanced binding to $G\alpha_s$ and $G\alpha_q$ compared with GIV. *A*, DAPLE binding to $G\alpha_s$ or $G\alpha_q$, but not to $G\alpha_{i3}$, is reduced upon replacing its GBA motif with that of GIV. *Upper panel*, diagram depicting the alignment of the GBA motifs of DAPLE and GIV and the sequence of the GBA motif of the DAPLE/GIV GBA chimera 1 (*ch1*) containing GIV's GBA motif residues (*red*) grafted into DAPLE's sequence (*black*). *Lower panel*, purified His– $G\alpha_{i3}$, His– $G\alpha_s$, or His– $G\alpha_q^*$ was incubated with GST, GST–DAPLE (WT or ch1), or GST–GIV immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. The *vertical dotted lines* indicate that the images were assembled by splicing lanes from the same experiment and membrane. *B*, mapping of residues involved in the differential G-protein selectivity of DAPLE *versus* GIV. *Upper panel*, sequences of DAPLE/GIV GBA chimeras (ch1–7, M1669V, and S1666G), with the GIV residues replaced in DAPLE indicated in *red. Lower panel*, purified His– $G\alpha_{i3}$, His– $G\alpha_s$, or His– $G\alpha_q^*$ was incubated with GST or GST–DAPLE (WT or mutants) immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. *C*, validation of the effects of DAPLE F1675A (FA) and M1669V (MV) mutations on binding to different G proteins using a shorter DAPLE-purified protein. *Upper panel*, diagram depicting the GST–DAPLE (short) construct used in this panel along with a diagram of the previously used GST–DAPLE construct. *Lower panel*, purified His– $G\alpha_{i3}$, His– $G\alpha_q^*$ was incubated with GST or GST–DAPLE (WT or mutants) immobilized on GSH-agarose beads. Bead-bound proteins were detected by Ponceau S staining or IB as indicated. All results presented in this figure are representative of at least three independent experiments ($n \ge 3$).

accelerate nucleotide exchange (55). Consistent with the results obtained with the constitutively-active mutants, loading of the G proteins with GDP·AlF $_4^-$ and/or GTP γ S markedly diminished binding to DAPLE compared with the GDP-loaded conditions (Fig. 2, B and D). Taken together, these results show that the interaction of DAPLE with $G\alpha_s$ or $G\alpha_q$ is G-protein state-dependent, having a marked preference for the inactive state. This feature resembles the previously characterized binding properties of GBA motifs to $G\alpha_s$ proteins (32, 33, 37–40).

Identification of a single amino acid in DAPLE that favors its binding to $G\alpha_s$ and $G\alpha_a$

Our results so far indicate that the GBA motif of DAPLE interacts with $G\alpha_s$ or $G\alpha_q$ through a mechanism that resembles binding of $G\alpha_i$ to previously described GBA motifs (*i.e.* direct binding to inactive $G\alpha$ subunits). Next, we set out to dissect the molecular determinants within the GBA motif of DAPLE that enable specific binding to $G\alpha_s$ and $G\alpha_q$. We reasoned that this specificity would be encoded in the GBA motif itself and that certain amino acids within the GBA motif of DAPLE that are different in the GBA motif GIV might confer specificity in G-protein binding. To test this hypothesis, we constructed a chimera of DAPLE in which 26 amino acids of the GBA motif of DAPLE were replaced by the corresponding amino acids from the GBA motif of GIV (GST-DAPLE ch1). We found that DAPLE ch1, much like GIV, fails to bind $G\alpha_s$ and $G\alpha_g$ but

retains binding to $G\alpha_{i3}$ (Fig. 3A). This indicates that amino acids within the GBA motif of DAPLE different from those in the GBA motif of GIV are required for its improved binding to $G\alpha_s$ and $G\alpha_g$. To map which one(s) of the differing amino acids is responsible for the observed difference in binding, we constructed six additional DAPLE/GIV GBA chimeras (GST-DAPLE ch2 to ch7) (Fig. 3B). These chimeras allowed us to reduce the number of amino acids that might confer increased binding to $G\alpha_s$ and $G\alpha_q$ to only two (Fig. 3*B*). Mutating each one of those two amino acids to the corresponding amino acid in GIV revealed that only M1669V reduces binding of DAPLE to $G\alpha_s$ and $G\alpha_g$ while still binding well to $G\alpha_{i3}$ (Fig. 3*B*). These results indicate that one single amino acid of the GBA motif of DAPLE, the methionine in position 1669, confers increased binding to $G\alpha_s$ and $G\alpha_g$ and that its mutation to a valine impairs these interactions.

In the course of our experiments, we observed that the different GST–DAPLE chimeras showed variable patterns of degradation products, which we reasoned might affect the validity of our conclusion that the M1669V mutation specifically impairs the interaction of DAPLE with $G\alpha_s$ or $G\alpha_q$. To address this issue, we constructed a shorter fragment of DAPLE fused to GST that we named GST–DAPLE (short), which we anticipated to show less degradation. We introduced in this construct the mutation F1675A that abrogates binding to all $G\alpha$ subunits,

and the mutation M1669V, which, based on the results showed above, should disrupt binding to $G\alpha_s$ and $G\alpha_g$ but not to $G\alpha_{i3}$. As expected, we found that the degradation products in GST-DAPLE (short) were greatly reduced and that the integrity of WT and mutant purified proteins were essentially the same. Moreover, we found that F1675A abolishes binding of DAPLE to all $G\alpha$ subunits tested, whereas M1669V only disrupts binding to $G\alpha_s$ and $G\alpha_g$ but not to $G\alpha_{i3}$ (Fig. 3C). These data confirm our results obtained with the longer fragment of DAPLE and indicate a key role for Met-1669 in mediating the binding to $G\alpha_s$ and $G\alpha_a$.

Met-1669 in DAPLE is highly conserved in evolution, as is the corresponding Val-1679 in GIV (Fig. S2A). Next, we asked whether mutation of GIV Val-1679 to the corresponding methionine found in DAPLE would be sufficient to enhance its binding to $G\alpha_s$ and $G\alpha_q$. We found that this is the case because GST-GIV V1679M bound to $G\alpha_s$ and to $G\alpha_g$ more efficiently than GST–GIV WT, whereas binding to $G\alpha_{i3}$ was not significantly affected (Fig. S2). These observations strengthen the conclusion that Met-1669 in DAPLE is the key determinant that allows efficient binding to $G\alpha_s$ and $G\alpha_g$.

Full-length DAPLE M1669V mutant displays impaired binding to $G\alpha_s$ and $G\alpha_q$ but not to $G\alpha_{i3}$

To determine whether the different specificity of DAPLE WT, F1675A, or M1669V for binding to $G\alpha$ observed in *in vitro* binding experiments with truncated proteins also occurs in the context of the full-length protein, we carried out co-IP experiments. FLAG-tagged $G\alpha_{i3}$, $G\alpha_{s}$, or $G\alpha_{g}$ was co-expressed with DAPLE WT, F1675A, or M1669V in HEK293T cells, and lysates were subjected to IP with FLAG antibodies. We found that DAPLE M1669V shows decreased binding to $G\alpha_s$ or $G\alpha_g$ but not to $G\alpha_{i3}$ (Fig. 4A), whereas DAPLE F1675A displays diminished binding to all three G proteins (i.e. $G\alpha_{i3}$, $G\alpha_s$, or $G\alpha_g$). These findings are in good agreement with our *in vitro* binding experiments, and they indicate that Met-1669 is required for binding of full-length DAPLE to $G\alpha_s$ or $G\alpha_g$ but is largely dispensable for interacting with $G\alpha_{i3}$.

To gain further insights into the structural basis for the role of Met-1669 in DAPLE and Val-1679 in the corresponding position of GIV in determining their different G-protein specificity, we leveraged the recently elucidated atomic resolution structure of the $G\alpha_{i3}$ /GIV GBA motif complex (35). We generated a homology model of the $G\alpha_{i3}/DAPLE$ GBA motif complex and compared the spatial localization of Met-1669 in DAPLE with that of Val-1679 in GIV (Fig. 4B). We observed that both GIV Val-1679 and DAPLE Met-1669 are largely solvent-exposed and do not make direct contact with $G\alpha_{i3}$ (Fig. 4B). This is very consistent with our results above showing that the DAPLE- $G\alpha_{i3}$ interaction tolerates well the replacement of Met-1669 in DAPLE by valine as determined in protein-protein-binding experiments above, and with previously published evidence showing that the GIV-G α_{i3} interaction tolerates well the replacement of Val-1679 in GIV by almost any other amino acid as determined in peptide array binding experiments (34). Overall, these observations provide a reasonable structural explanation for the neutral role of the GIV V1679/DAPLE Met-1669 position in determining binding to $G\alpha_{i3}$. Although we lack

a reliable template to generate high-confidence homology models of DAPLE's GBA motif in complex with $G\alpha_s$ or $G\alpha_g$, we propose that the larger side chain of DAPLE Met-1669 compared with GIV Val-1679 might allow for additional molecular contacts with $G\alpha_s$ and $G\alpha_g$ and thereby account for its ability to bind $G\alpha_s$ and $G\alpha_g$ better than GIV (Fig. 4C). Such contacts would be disrupted upon mutation of Met-1669 to valine, which in turn would have no effect on $G\alpha_{i3}$ binding (Fig. 4*C*).

Peptide derived from the GBA motif of DAPLE recapitulates its GEF activity on $G\alpha_{i3}$

Next, we set out to evaluate the consequences of DAPLE binding to $G\alpha_s$ and $G\alpha_g$ on G-protein activity. We reasoned that a peptide derived from the GBA motif of DAPLE could be used for this purpose, as previous observations suggest that GBA peptides recapitulate well the properties of their cognate proteins in that they regulate $G\alpha_i$ proteins (34, 35). We fully validated these previous observations by analyzing the dose-dependent effects of a 34-mer GBA peptide from DAPLE in two assays that monitor nucleotide exchange on $G\alpha_i$: steady-state GTPase and GTPyS binding assays (56). Briefly, steady-state GTPase activity is a good proxy of nucleotide exchange rates for $G\alpha_i$ because hydrolysis is 1–2 orders of magnitude faster than nucleotide exchange, whereas GTP γ S binding is a more direct readout of exchange (57). As expected, we found that the DAPLE GBA peptide, but not a control peptide (see "Experimental procedures"), increased the steady-state GTPase activity (Fig. S3, A and B) and GTP γ S binding (Fig. S3C) of $G\alpha_{i3}$ in a dose-dependent manner. The amplitude of the DAPLE GBA peptide effects (\sim 2–4-fold increases) and the corresponding EC₅₀ values (\sim 5–10 μ M) are similar to those previously reported for a very similar peptide from GIV whose molecular mechanism of action on $G\alpha_i$ has been extensively validated (34, 35). We conclude that the GBA peptide of DAPLE recapitulates the GEF activity of DAPLE protein *in vitro*.

DAPLE inhibits nucleotide exchange on $G\alpha_{\varsigma}$

As for $G\alpha_{i3}$, we determined the effects of DAPLE on $G\alpha_{s}$ nucleotide exchange using both steady-state GTPase assays and GTP_{\gamma}S-binding assays. We found that DAPLE GBA peptide decreases $G\alpha_s$ steady-state GTPase activity \sim 40% (Fig. 5, A and *B*), which was in agreement with an \sim 50% decrease in the rate of GTP γ S binding (Fig. 5C). The half-maximal inhibitory concentration (IC₅₀) was \sim 3 μ M in both assays (Fig. 5, B and D). These results indicate that DAPLE GBA peptide inhibits nucleotide exchange on $G\alpha_s$ and that the potency of this inhibition is similar to or higher than the potency it has for activating $G\alpha_{i3}$ (\sim 5–10 μ M, Fig. S3).

To rule out that the observed effect on $G\alpha_s$ activity was related to the use of an isolated GBA motif out of protein context, we performed additional experiments with a larger fragment of DAPLE. For this, we used a purified protein consisting of the GBA-containing C-terminal region of DAPLE (DAPLE-CT, aa 1650 – 2028), which has been previously validated to preserve the G-protein regulatory functions of DAPLE on $G\alpha_{i3}$ (39). We found that purified DAPLE-CT slows down GTP γ S binding to $G\alpha_s$ to an extent similar to that observed with



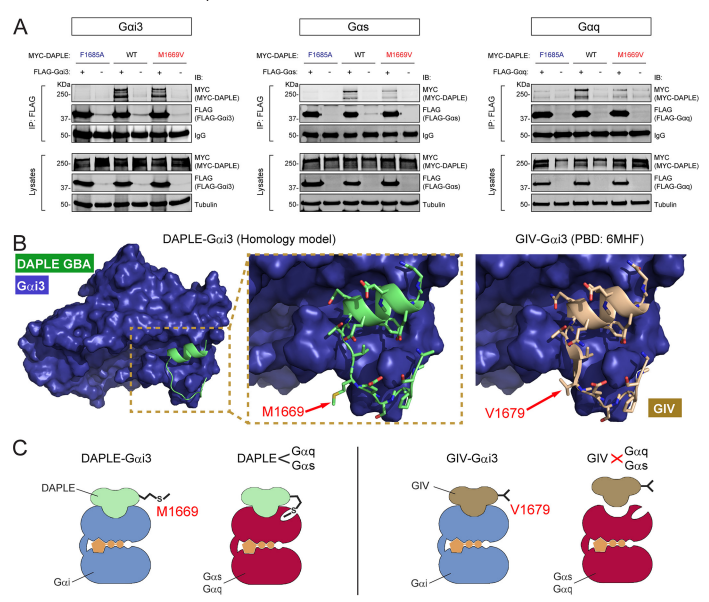


Figure 4. M1669V mutation in full-length DAPLE disrupts binding to $G\alpha_s$ or $G\alpha_q$ but not to $G\alpha_{13}$. A, co-immunoprecipitation experiments comparing the effect of DAPLE M1669V and F1675A mutations on G-protein binding, which show that the former disrupts binding to $G\alpha_s$ and $G\alpha_{qr}$ but not to $G\alpha_{13}$, whereas the latter disrupts binding to all G proteins tested. Lysates of HEK293T cells co-expressing full-length MYC-DAPLE (WT or mutants) with the indicated FLAG-tagged G proteins (or no tagged G protein as negative control) were subjected to IP with a FLAG antibody, and bound proteins were detected by IB as indicated. The *lower immunoblot panels* (*Lysates*) correspond to aliquots of the starting material used for IPs shown in the *upper panels* (*IP: FLAG*). One representative experiment of four is shown for $G\alpha_s$ and $G\alpha_q$ (n=4), or one representative experiment of two is shown for $G\alpha_{13}$ (n=2). B, comparison of DAPLE Met-1669 and GIVVal-1679 in the context of their respective $G\alpha_{13}$ /GBA motif complex structures. *Left panel*, homology model of DAPLE GBA motif (*green, ribbon* representation) in complex with $G\alpha_{13}$ (*blue*, space-filling representation) was generated using the X-ray crystal structure of the $G\alpha_{13}$ /GIV GBA motif complex (PDB code 6MHF). The area of the $G\alpha_{13}$ /DAPLE structure model within the *dotted box* is shown enlarged in the *middle panel* to illustrate that Met-1669 is largely solvent-exposed. *Right panel*, detail of the structure of $G\alpha_{13}$ in complex with GIV GBA motif (*brown*) showing that Val-1669 is also largely solvent-exposed. *C*, proposed model for the structural basis of DAPLE's G-protein selectivity. Much like GIV Val-1679, DAPLE Met-1669 does not make direct contact with $G\alpha_{13}$. In contrast, DAPLE Met-1669 is required for binding to $G\alpha_s$ or $G\alpha_q$, suggesting that it makes a contact with these proteins that is not allowed by the shorter chain of the valine located in the corresponding position in GIV.

DAPLE GBA peptide (\sim 50%, Fig. 5*E*). Moreover, the inhibitory effect of DAPLE-CT on $G\alpha_s$ was blunted by the F1675A mutation that disrupts *G*-protein binding (Fig. 5*F*), confirming that this effect is mediated by the GBA motif of DAPLE. Altogether, these results indicate that the GBA motif of DAPLE can *inhibit* nucleotide-exchange activity of $G\alpha_s$ as potently as it activates $G\alpha_i$ and that this occurs in the absence of any phosphomodification because the experiments were performed with bacterially-expressed proteins or synthetic peptides.

DAPLE inhibits nucleotide exchange on $G\alpha_a$

Next, we investigated the effect of DAPLE on the activity of $G\alpha_{\rm q}$. Instead of using the bacterially-expressed $G\alpha_{\rm q}^*$ chimera utilized in our protein-binding experiments described above, we used $G\alpha_{\rm q}$ purified from Sf9 insect cells because its enzymatic properties have been more thoroughly characterized (55, 58), including its modulation by various regulators (17, 59). $G\alpha_{\rm q}$ differs markedly from $G\alpha_{\rm i}$ or $G\alpha_{\rm s}$ in that its spontaneous exchange of nucleotide is very slow (55, 58). Thus, steady-state

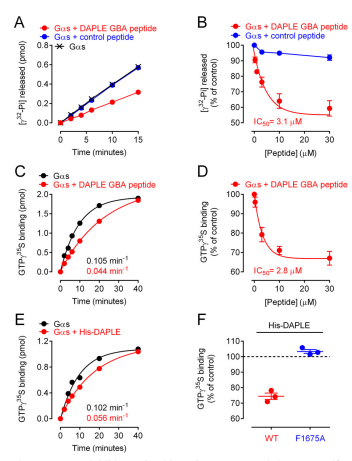


Figure 5. DAPLE inhibits nucleotide exchange on $G\alpha_s$ via its GBA motif. A and B, DAPLE GBA peptide decreases the steady-state GTPase activity of $G\alpha_s$. A representative time course of the steady-state GTPase activity of His-G α_s alone (black), in the presence of DAPLE GBA peptide (30 μM, red), or control peptide (30 μ M, blue) is shown in A, and quantification of the dose-dependent effect of the peptides is shown in B (mean \pm S.E., n = 3). Results are presented as raw production of free $[^{32}P]P_i$ (pmol) in A or percent change relative to the production of free [32 P]P, by G $\alpha_{\rm s}$ alone at 10 min (% of control) in *B.* Average IC₅₀ value was determined as described under "Experimental procedures." and D, DAPLE GBA peptide decreases the rate of GTP γ S binding to G α_s . A representative time course of [35 S]GTP γ S binding to His–G $\alpha_{\rm s}$ in the absence (black) or presence of DAPLE GBA peptide (30 μ M, red) is shown in C, and quantification of the dose-dependent effect of the DAPLE GBA peptide is shown in D (mean \pm S.E., n=3). Results are presented as raw [35 S]GTP γ S binding (picomoles) in C or percent change relative to [35 S]GTP γ S binding to $G\alpha_s$ alone at 10 min (% of control) in D. Rate constants and average IC_{50} values were determined as described under "Experimental procedures," purified DAPLE WT (amino acids 1650-2028), but not F1675A mutant, decreases GTP γ S binding to G α_s . A representative time course of [35 S]GTP γ S binding to His–G α_s in the absence (*black*) or presence of purified His–DAPLE (9 μ M, red) is shown in E, and quantification of the effect of His–DAPLE WT (3.3 μ M, red) compared with His–DAPLE F1675A (3.3 μ M, blue) is shown in D (mean \pm S.E., n= 3). Results are presented as raw [35 S]GTP γ S binding (picomoles) in E or percent change relative to [35 S]GTP γ S binding to G α_s alone at 10 min (% of control) in F. Rate constants determined as described under "Experimental procedures."

GTPase activity is not a good proxy for nucleotide exchange, so we exclusively measured GTP yS binding to assess the effects of DAPLE on nucleotide exchange. Moreover, we supplemented the assay buffer with 0.2 M ammonium sulfate, as described previously (55), to enhance nucleotide exchange to a faster rate that is more tractable experimentally. Under these conditions, we found that DAPLE GBA peptide slowed down GTP \(\gamma \) binding ${\sim}40\%$, with an IC $_{50}$ of ${\sim}3~\mu\mathrm{M}$ (Fig. 6, A and B). This effect was recapitulated by DAPLE-CT WT but not the G-protein binding-deficient F1675A mutant (Fig. 6C), indicating that $G\alpha_{\alpha}$

inhibition is a bona fide action of the GBA motif of DAPLE. We ruled out that the observed inhibition was a consequence of performing the experiments under artificially accelerated nucleotide-exchange conditions by performing analogous experiments with a buffer not supplemented with ammonium sulfate (Fig. S3). As expected, the rate of nucleotide exchange was much slower, but the overall effects of DAPLE GBA peptide (Fig. S4, A and B) and DAPLE-CT (Fig. S4C) were analogous, i.e. DAPLE inhibited nucleotide exchange with similar potency and efficacy. Taken together, these results show that DAPLE *inhibits* nucleotide exchange on $G\alpha_{\alpha}$ via its GBA motif.

Discussion

DAPLE belongs to a family of cytoplasmic G-protein regulators that are defined by the presence of a GBA motif. Although proteins with a GBA motif have been previously found to preferentially bind to $G\alpha$ subunits of G proteins of the $G_{i/o}$ family over $G\alpha$ subunits of other G-protein families, the main discovery of the work presented here is that DAPLE binds to $G\alpha_s$ (G_s family) and to $G\alpha_q$ ($G_{q/11}$ family). Moreover, although GBA motifs have been characterized by their ability to exert GEF activity on $G\alpha_i$ subunits, here we found that DAPLE exerts GDI activity on $G\alpha_s$ and $G\alpha_g$. Previously reported evidence indicates that GIV, another protein with a GBA motif, can bind to $G\alpha_s$ and exert GDI activity on it (51). However, this G-protein regulatory function on $G\alpha_{\epsilon}$ appeared only after sequential phosphorylation of two sites that flank the GBA motif, which also precluded $G\alpha_i$ binding and activation by GIV (51). This implies that GIV can work as either a GEF or a GDI depending on a switch of G-protein-binding preference determined by posttranslational phosphorylation. In contrast, this work shows that, in the absence of post-translational phosphorylation, DAPLE binds to $G\alpha_s$ or $G\alpha_q$ and exerts GDI activity on them, while still retaining the ability to bind and regulate $G\alpha_i$. This implies that DAPLE can work as a GEF for $G\alpha_i$ and a GDI for $G\alpha_s$ and $G\alpha_q$ (Fig. 7). To our knowledge, this is also the first description of a protein, other than $G\beta\gamma$ dimers, with GDI activity toward $G\alpha$ subunits of the $G_{\alpha/11}$ family. These findings expand on previous observations with synthetic peptides similar to GBA motifs that were shown to bind and regulate $G\alpha$ proteins. For example, the synthetic peptide KB-752, based on which was the first GBA motif in GIV identified by similarity (32), was initially shown to be a GEF for $G\alpha_i$ (60) but later to also be a GDI for $G\alpha_s$ (61). This dual-specificity and bi-functional regulatory action on G proteins was also described for another GBA-like synthetic peptide, namely GSP, identified independently (62). Thus, although all other proteins with a GBA motif described to date have been shown to have GEF activity for $G\alpha_i$ in vitro (32, 33, 38-40), it still remains to be elucidated whether they can also work as GDIs depending on their post-translational modification status and/or the nature of their G-protein substrate.

Here, we also define the similarities and differences of the interaction between DAPLE and $G\alpha_s$ or $G\alpha_q$ compared with $G\alpha_i$. As reported previously for the interaction of DAPLE with $G\alpha_i$ (and with GBA motifs of other proteins), DAPLE also binds preferentially to the inactive conformation of $G\alpha_s$ or $G\alpha_g$. This interaction is ablated upon mutation of a hydrophobic amino



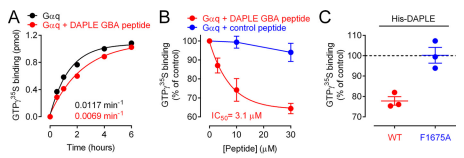


Figure 6. DAPLE inhibits nucleotide exchange on $G\alpha_q$ via its GBA motif. A and B, DAPLE GBA peptide decreases the rate of GTPγS binding to $G\alpha_q$. A representative time course of [35 S]GTPγS binding to $G\alpha_q$ in the absence (black) or presence of DAPLE GBA peptide ($30 \mu M$, red) using a buffer that contains 0.2 M (NH₄) $_2$ SO₄ is shown in A, and quantification of the dose-dependent effect of DAPLE GBA peptide (red) or control peptide (blue) in a buffer that contains 0.2 M (NH₄) $_2$ SO₄ is shown in B (mean \pm S.E., n = 3). Results are presented as raw [35 S]GTPγS binding (picomoles) in A or percent change relative to [35 S]GTPγS binding to $G\alpha_q$ alone at 45 min (% of control) in B. Rate constants and average IC $_{50}$ values were determined as described under "Experimental procedures." C, purified DAPLE WT (amino acids 1650–2028), but not F1675A mutant, decreases GTPγS binding to $G\alpha_q$. Quantification of the effect of His–DAPLE WT (3.3 μ M, red) compared with His–DAPLE F1675A (3.3 μ M, blue) in a buffer that contains 0.2 M (NH₄) $_2$ SO₄ is presented as percent change relative to [35 S]GTPγS binding to $G\alpha_q$ alone at 45 min (% of control, mean \pm S.E., n = 3).

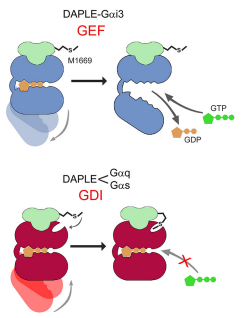


Figure 7. Proposed model. *Top,* DAPLE (*green*) binds to $G\alpha_i$ (*blue*) to stabilize a G-protein conformation that favors nucleotide exchange. DAPLE Met-1669 is not required for efficient binding to $G\alpha_i$. *Bottom,* Met-1669 in DAPLE allows its efficient physical engagement to $G\alpha_s$ and $G\alpha_q$ (*red*), which in turn stabilizes a G-protein conformation that prevents nucleotide exchange.

acid (Phe-1675) that is highly conserved across GBA motifs of different proteins and is also essential for $G\alpha_i$ binding (37, 40, 47). This suggests that the overall binding mode of DAPLE's GBA motif on $G\alpha_s$ or $G\alpha_q$ probably resembles that observed for $G\alpha_i$, *i.e.* docking onto a hydrophobic cleft formed by the switch II and α 3 helix of $G\alpha$ that becomes inaccessible upon G-protein activation due to conformational rearrangement of switch II (32, 37). Regarding the differences between DAPLE binding to $G\alpha_s/G\alpha_a$ versus $G\alpha_i$, we identified here a single amino acid in DAPLE (Met-1669) that is required for binding to $G\alpha_s$ and $G\alpha_g$ but not to $G\alpha_i$. Although the existence of this additional molecular contact between DAPLE and $G\alpha_s/G\alpha_a$ correlates with the ability to exert GDI activity, we can only speculate about the mechanism by which this additional contact might inhibit nucleotide exchange on $G\alpha$ based on previously described observations in the literature. Essentially, there are two mechanisms of action described for proteins with GDI activity. One

is that mediated by GoLoco motifs, which prevent nucleotide exchange by directly engaging GDP and securing it in the nucleotide-binding pocket through an "arginine finger" (18, 19). The other mechanism is that mediated by $G\beta\gamma$ subunits, which, instead of making direct contact with GDP, bind to the Ras-like domain of $G\alpha$ and allosterically stabilize GDP in the nucleotide-binding pocket (4, 63, 64). For the GDI action of DAPLE, we favor a mechanism akin to the latter rather than to the former. This is because, based on the discussion above, it is unlikely that the overall docking pose of DAPLE on $G\alpha_s/G\alpha_q$ differs markedly from that observed for $G\alpha_i$, so the GBA motif would not make direct contact with the nucleotide. Instead, the additional G-protein contact established through Met-1669 might favor a conformation of $G\alpha$ that is less prone to exchange nucleotide (Fig. 7). Elucidation of atomic resolution structures of DAPLE's GBA motif in complex with $G\alpha_s$ and/or $G\alpha_g$ would be required to shed light on this matter.

Finding that DAPLE has GDI activity toward $G\alpha_s/G\alpha_a$ sets a new framework to discover and/or understand biological functions of DAPLE's GBA motif. From a traditional viewpoint of GPCR signaling, GDI activity exerted on $G\alpha_s$ and $G\alpha_a$ could be expected to suppress their ability to engage and modulate their respective effectors, such as adenylyl cyclase for $G\alpha_s$ or PLC- β and RhoGEFs for $G\alpha_{g}$. However, preliminary evidence suggests that DAPLE's GBA motif does not influence GPCR-mediated modulation of some of these effector pathways.5 This is not entirely surprising because it is difficult to predict the context (e.g. specific GPCR) in which DAPLE might engage G proteins to regulate them. Previous reports with other GDIs, like the GoLoco motif-containing protein AGS3, have also found that GPCR/G-protein effector pathways are not necessarily impacted as one would predict based on their putative ability to prevent $G\alpha$ -effector signaling (65). Instead, GoLoco motif-containing GDIs have been shown to be engaged into alternative modes of G-protein signaling (66, 67). One of them consists of forming complexes with $G\alpha$ subunits that do not mediate signaling through GPCRs but are involved in the control of cell division in metazoans (68-71). In addition, the GoLoco motif containing GDIs was originally discovered in a yeast genetic

⁵ A. Marivin, M. Maziarz, and M. Garcia-Marcos, unpublished observations.

screen based on their ability to enhance $G\beta\gamma$ -dependent signaling (72). It has been subsequently proposed that, while having inhibitory GDI activity on $G\alpha$ subunits, GoLoco motifs promote $G\beta\gamma$ -dependent signaling by favoring $G\alpha$ - $G\beta\gamma$ dissociation (73, 74). It is possible that the GDI activity of DAPLE on $G\alpha_s/G\alpha_g$ results in a signaling mechanism in cells similar to that proposed for GoLoco motif-containing GDIs, as it has been reported that the physiological functions of DAPLE's GBA motif during embryonic development are $G\beta\gamma$ -dependent (41), and because it has been previously shown that GBA motifs can displace $G\beta\gamma$ subunits from $G\alpha$ – $G\beta\gamma$ complexes (32, 39). It is also important to take into account that DAPLE is a modular multifunctional protein and that such modularity might have a significant impact on the context in which DAPLE's GDI function operates. Most notably, DAPLE contains a PDZ-binding motif (PBM) that is required for its localization and function at apical cell-cell junctions (41, 49). Without this PBM, DAPLE's GBA motif cannot promote actomyosin contractility and the subsequent effects on embryonic morphogenesis (41). Thus, the functional outcomes of DAPLE's GDI activity might be spatially restricted to apical cell junctions, where different G-protein subtypes can localize (75–78). Further investigation will be required to elucidate the biological consequences of DAPLE's GDI activity on $G\alpha_s$ and/or $G\alpha_g$.

In summary, the findings reported here reveal that a protein with a GBA motif, namely DAPLE, can promiscuously bind to $G\alpha$ subunits of different subfamilies and that, depending on the G protein subtype, it exerts opposing actions on nucleotide exchange: it inhibits nucleotide exchange on $G\alpha_s$ and $G\alpha_{ab}$ while it accelerates it on $G\alpha_i$. These findings highlight the versatility of GBA motifs in modulating the activity of G proteins.

Experimental procedures

Reagents and antibodies

Unless otherwise indicated, all chemical reagents were obtained from Sigma or Thermo Fisher Scientific. Escherichia $coli \, DH5\alpha$ strain was purchased from New England Biolabs and the BL21(DE3) strain from Life Technologies, Inc. PfuUltra DNA polymerase was purchased from Agilent. $[\gamma^{-32}P]GTP$ and [35S]GTPyS were from PerkinElmer Life Sciences. Mouse monoclonal antibodies raised against α -tubulin (T6074), FLAG tag (F1804), or His tag (H1029) were from Sigma. Mouse mAb raised against hemagglutinin (HA) tag (clone 12CA5, catalog no. 11583816001) was obtained from Roche Applied Science. Mouse mAb raised against MYC tag (9B11, catalog no. 2276) was from Cell Signaling. Rabbit polyclonal antibodies raised against G $\alpha_{\rm s}$ (C-18, sc-383) or G $\alpha_{\rm q}$ (E-17, sc-393) were purchased from Santa Cruz Biotechnology. Goat anti-rabbit Alexa Fluor 680 (A21077) and goat anti-mouse IRDve 800 (catalog no. 926-32210) secondary antibodies were from Life Technologies, Inc., and LI-COR, respectively.

Plasmids

Plasmids for expression of GST-DAPLE (human aa 1650 – 2028, pGEX-4T-DAPLE-CT) and GST-GIV (human aa 1671-1755, pGEX-4T-GIV-CT) in bacteria were described previously (39). GST-DAPLE (short) (human aa 1650-1745) was generated by introduction of a stop codon in position 1746 of pGEX-4T-DAPLE-CT using site-directed mutagenesis. The plasmid for expression of His-DAPLE (human, aa 1650 – 2028, pET28b-DAPLE) in bacteria was described previously (39). Plasmids for expression of FLAG– $G\alpha_{i3}$ (rat, p3XFLAG-CMV10-G α_{i3} , N-terminal 3XFLAG tag), $G\alpha_{i3}$ -FLAG (rat, p3XFLAG-CMV14-G α_{i3} , C-terminal 3XFLAG tag), or $G\alpha_{i3}$ (human, pcDNA3.1(+)- $G\alpha_s$) in mammalian cells were described previously (32, 52). Plasmids for the expression of $G\alpha_q$ -HA (mouse, pcDNA3- $G\alpha_q$ -HA, internally tagged) or $G\alpha_{12}$ –MYC (mouse, pcDNA3.1- $G\alpha$ 12–MYC, internally tagged) in mammalian cells were kindly provided by P. Wedegaertner (Thomas Jefferson University) (79) and T. Meigs (University of North Carolina, Asheville) (80), respectively. Plasmid for the expression of His- $G\alpha_{i3}$ (rat, pET28b- $G\alpha_{i3}$) was described previously (32). Plasmids for the expression of His- $G\alpha_s$ (bovine, pHis6- $G\alpha_s$) and His- $G\alpha_q^*$ (mouse, pET28a- $G\alpha_{a}^{*}$) in bacteria were kindly provided by N. Artemyev (University of Iowa) and S. Sprang (University of Montana), respectively (53, 81). Plasmids for expression of $G\alpha_a$, $G\beta1$ –His, and $G\gamma 2$ –His in Sf9 insect cells were described previously (59). Plasmid for the expression of full-length MYC-DAPLE (human, pCS2-6XMYC-DAPLE) in mammalian cells was described previously (41). All point mutations, including DAPLE/GIV GBA chimeras, were generated by site-directed mutagenesis following the manufacturer's instructions (Quik-Change II, Agilent).

Expression and purification of proteins

GST, GST-DAPLE, GST-GIV, GST-DAPLE (short), and His- $G\alpha_{i3}$ proteins were purified from bacteria as described previously (32, 39). Briefly, proteins were expressed in BL21(DE3) E. coli (Life Technologies, Inc.) transformed with the corresponding plasmids by overnight induction at 23 °C with 1 mm isopropyl β -D-1-thio-galactopyranoside (IPTG) when the OD_{600} reached \sim 0.7. Bacteria were pelleted and resuspended at 4 °C in lysis buffer: 50 mм NaH₂PO₄, pH 7.4, 300 mm NaCl, 10 mm imidazole, and 1% (v/v) Triton X-100 supplemented with protease inhibitor mixture (1 µM leupeptin, 2.5 μM pepstatin, 0.2 μM aprotinin, and 1 mM PMSF). For $G\alpha_{i3}$ this buffer was also supplemented with 25 μ M GDP and 5 mM MgCl₂. After sonication (30-s bursts, four times), lysates were cleared by centrifugation at 12,000 \times g for 30 min at 4 °C. The supernatant was used for affinity purification on HisPur Cobalt or GSH-agarose resins (Thermo Fisher Scientific) for 90 min at 4 °C with rotation. Resins were washed four times with lysis buffer, and proteins were eluted with lysis buffer supplemented with 250 mm imidazole or with 50 mm Tris-HCl, pH 8, 100 mm NaCl, and 30 mm reduced GSH for HisPur Cobalt and GSHagarose resins, respectively. The eluted fractions were dialyzed overnight at 4 °C against PBS, except for $G\alpha_{i3}$, which was buffer exchanged into 20 mm Tris-HCl, pH 7.4, 20 mm NaCl, 1 mm MgCl₂, 1 mm, DTT, 10 μM GDP, and 5% (v/v) glycerol using a HiTrap desalting column (GE Healthcare). All protein samples were aliquoted and stored at -80 °C.

His- $G\alpha_s$ was purified from bacteria as described previously (82). Briefly, His-G α_s was expressed in BL21(DE3) *E. coli* transformed with the corresponding plasmid by overnight induction at 23 °C with 0.1 mm IPTG when OD_{600} reached \sim 0.5. Bacteria



were pelleted and resuspended at 4 °C in lysis buffer (50 mm Tris-HCl, pH 8.0, 50 mm NaCl, 5 mm MgCl₂, 50 μ m GDP, and 5 mm β -mercaptoethanol) supplemented with protease inhibitor mixture (1 μ M leupeptin, 2.5 μ M pepstatin, 0.2 μ M aprotinin, and 1 mM PMSF). After sonication (30-s bursts, four times), the lysate was cleared by centrifugation at 12,000 \times g for 30 min at 4°C. The supernatant was adjusted to 500 mм NaCl and 20 mм imidazole before affinity purification by incubation with nickelnitrilotriacetic acid resin (Qiagen) for 90 min at 4 °C. Resin was washed four times with lysis buffer, and protein was eluted with lysis buffer supplemented with 100 mM imidazole. The eluted fraction was adjusted to 50 mm Tris-HCl, pH 8.0, 50 mm NaCl, 5 mm MgCl₂ using a protein concentrator with a 10-kDa cutoff (Pierce) before loading onto a HiTrap Q HP column (GE Healthcare) connected to an ÄKTA FPLC. Proteins were eluted by applying a 50 – 500 mm NaCl gradient, and fractions containing His-G α_s were pooled and supplemented with 10 μ M GDP and 5 mm β -mercaptoethanol before concentration in 50 mm Tris-HCl, pH 8.0, 150 mm NaCl, 5 mm MgCl₂, 5% (w/v) glycerol, 10 μ M GDP, and 5 mM β -mercaptoethanol, and storage at -80 °C.

His– $G\alpha_q^*$ was purified from bacteria as described previously (53). His- $G\alpha_{\alpha}^*$ was expressed in BL21(DE3) *E. coli* transformed with the corresponding plasmid by overnight induction at 23 °C with 1 mm IPTG when OD_{600} reached \sim 0.7. Bacteria were pelleted and then resuspended at 4 °C in lysis buffer: 20 mm HEPES, pH 8.0, 300 mm NaCl, 5 mm MgCl₂, 10 mm β -mercaptoethanol, 15 mm imidazole, 10% (v/v) glycerol, 50 μm GDP, 30 μΜ AlCl₃, and 10 mm NaF supplemented with protease inhibitor mixture (1 μM leupeptin, 2.5 μM pepstatin, 0.2 μM aprotinin, and 1 mM PMSF). After sonication (30-s bursts, four times), the lysate was cleared by centrifugation at 12,000 \times g for 30 min at 4 °C. The supernatant was used for affinity purification on His-Pur Cobalt resin (Pierce) for 90 min at 4 °C with rotation. The resin was washed four times with lysis buffer, and the protein was eluted with lysis buffer supplemented with 500 mm imidazole. The eluted fraction was supplemented with 100 mm EDTA and incubated on ice for 20 min before loading onto a Superdex 200 HR10/30 gel-filtration column using a running buffer consisting of 20 mm HEPES, pH 8.0, 100 mm NaCl, 1 mm $MgCl_2$, 2 mm DTT, 2% (v/v) glycerol, and 10 μ m GDP. Fractions containing His–G α_q^* were concentrated and stored at –80 °C. $G\alpha_{a}$ was purified from Sf9 insect cells as described previously (83).

In vitro protein-binding assays with GST-fused proteins

The following GST-fused proteins were immobilized on GSH-agarose beads for 90 min at room temperature in PBS (amounts per condition are indicated in parentheses): GST (30 μ g), GST-DAPLE (30 μ g), GST-GIV (30 μ g), and GST-DAPLE (short) (30 μ g). Beads were washed twice with PBS and resuspended in 300 μ l of binding buffer (50 mM Tris-HCl, pH 7.4, 100 mM NaCl, 0.4% (v/v) Nonidet P-40, 5 mM EDTA, 2 mM DTT) supplemented with 30 μ M GDP unless indicated otherwise.

For experiments using cell lysates as a source of soluble binding ligands, HEK293T cells (ATCC CRL-3216) were grown at 37 °C, 5% CO₂ in high-glucose Dulbecco's modified Eagle's

medium (DMEM) supplemented with 10% fetal bovine serum, 100 units/ml penicillin, 100 μ g/ml streptomycin, and 1% L-glutamine. Approximately two million HEK293T cells were seeded on 10-cm dishes and transfected the day after using the calcium phosphate method with plasmids encoding the following constructs (DNA amounts in parentheses): $G\alpha_{i3}$ -FLAG (3 μ g), $G\alpha_{s}$ $(3 \mu g)$, $G\alpha_q$ –HA $(6 \mu g)$, or $G\alpha_{12}$ –MYC $(3 \mu g)$. Cell medium was changed 6 h after transfection. Thirty two hours after transfection, cells were lysed at 4 °C with 700 µl of lysis buffer (20 mm HEPES, pH 7.2, 125 mm K(CH₃COO), 0.4% (v/v) Triton X-100, 1 mm DTT, 10 mm β-glycerophosphate, and 0.5 mm Na₃VO₄ supplemented with a protease inhibitor mixture (SigmaFAST, catalog no. S8830)). Cell lysates were cleared by centrifugation at $14,000 \times g$ for 10 min. For experiments using nucleotideloaded $G\alpha_s$, the lysates of HEK293T were incubated at 30 °C for 30 min with nucleotides as follows: 125 μM GDP (GDP condition), or 125 μM GDP, 125 μM AlCl₃, 10 mM NaF (GDP·AlF₄ condition), or 125 μ M GTP γ S (GTP γ S condition). For experiments using nucleotide-loaded $G\alpha_q$, the lysates were incubated at 4 °C for 30 min with nucleotides as follows: 30 μM GDP (GDP condition), or 30 μM GDP, 30 μM AlCl₃, 10 mM NaF (GDP·AlF₄

One hundred microliters (\sim 400 μ g) of cell lysate prepared as described above were added to the GST-fused proteins immobilized on resin beads and incubated 4 h at 4 °C with constant rotation. Beads were washed four times with 1 ml of wash buffer (4.3 mm Na₂HPO₄, 1.4 mm KH₂PO₄, pH 7.4, 137 mm NaCl, 2.7 mm KCl, 0.1% (v/v) Tween 20, 5 mm EDTA, 1 mm DTT) supplemented with 30 μ m GDP, except for experiments using G α loaded with GDP·AlF $_4^-$ or GTP γ S, in which the wash buffer was supplemented with 30 μ m GDP, 30 μ m AlCl₃, and 10 mm NaF, or with 30 μ m GTP γ S, respectively. Resin-bound proteins were eluted by boiling for 5 min in Laemmli sample buffer, and proteins were separated by SDS-PAGE and immunoblotted with antibodies as indicated under "Immunoblotting."

For experiments using purified proteins as source of soluble binding ligands, purified proteins were prepared as described under "Expression and purification of proteins." Aliquots of protein stored at $-80\,^{\circ}\mathrm{C}$ were quickly thawed and cleared by centrifugation at $14,\!000\times g$ for 2 min before addition to GST-fused proteins immobilized on GSH-agarose beads in 300 $\mu\mathrm{l}$ of binding buffer supplemented with 30 $\mu\mathrm{M}$ GDP. The amount of purified proteins used were as follows: 1.7 $\mu\mathrm{g}$ of His–G α_{i3} , 1.65 $\mu\mathrm{g}$ of His–G α_s , and 1.7 $\mu\mathrm{g}$ of His–G α_q^* . Tubes were incubated for 4 h at 4 °C with constant rotation, and washes and elution were performed as described above. Proteins were separated by SDS-PAGE and immunoblotted with antibodies as indicated under "Immunoblotting."

Immunoprecipitation

Approximately three million HEK293T cells were seeded in 10-cm dishes and transfected the day after using polyethyleneimine (PEI; Polysciences, Inc.; catalog no. 23966, 1 mg/ml solution reconstituted in water). For each 10-cm dish, a total of 9 μ g of DNA were transfected, consisting of 3 μ g of plasmids encoding FLAG–G α_{i3} , FLAG–G α_{s} , FLAG–G α_{q} , or an empty vector, along with 6 μ g of MYC–DAPLE (WT or mutants). Plasmid DNA was added to 500 μ l of DMEM and immediately mixed

with PEI reagent (DNA/PEI reagent ratio of 1:3) by vortexing for 2 s. Tubes were incubated at room temperature for 15 min before adding to cells, and media were changed 6 h later. Twenty four hours after transfection, cells were lysed on ice with 750 µl of lysis buffer (20 mm HEPES, pH 7.2, 5 mm Mg(CH₃COO)₂, 125 mm K(CH₃COO), 0.4% (v/v) Triton X-100, 1 mm DTT, 10 mm β-glycerophosphate, 0.5 mm Na_3VO_4 , and 30 μ M GDP supplemented with a protease inhibitor mixture (Sigma S8830)) and cleared (14,000 \times g, 10 min). Cleared lysates were incubated with 2 µg of FLAG antibodies (Sigma F1804) for 2.5 h at 4 °C with constant rotation. Forty μl of an ~50% protein G-agarose beads suspension, pre-blocked with 5% (w/v) BSA in PBS for 2 h at room temperature, were added to the lysate/antibody mixture and incubated for 90 min at 4 °C. Beads were washed three times with wash buffer (4.3 mм Na₂HPO₄, 1.4 mм KH₂PO₄, pH 7.4, 137 mм NaCl, 2.7 mм KCl, 0.1% (v/v) Tween 20, 10 mm MgCl₂, 5 mm EDTA, 1 mm DTT, and 30 μ M GDP), and proteins were eluted by boiling in Laemmli sample buffer for 5 min. Proteins were separated by SDS-PAGE and immunoblotted with antibodies as indicated under "Immunoblotting."

Immunoblotting

Proteins were separated by SDS-PAGE and transferred to PVDF membranes, which were blocked with 5% (w/v) nonfat dry milk and sequentially incubated with primary and secondary antibodies. For protein-protein-binding experiments with GST-fused proteins, PVDF membranes were stained with Ponceau S and scanned before blocking. The primary antibodies used were the following: MYC (1:1000); His (1:2500); FLAG (1:2000); α -tubulin (1:2500); HA (1:1000); G α_s (1:500); and G α_a (1:500). The secondary antibodies were goat anti-rabbit Alexa Fluor 680 (1:10,000) and goat anti-mouse IRDye 800 (1:10,000). IR imaging of immunoblots was performed using an Odyssey IR Imaging System (Li-Cor Biosciences). Images were processed using ImageJ software (National Institutes of Health) and assembled for presentation using Photoshop and Illustrator softwares (Adobe).

Homology modeling

A model of $G\alpha_{i3}$ bound to the GBA motif of DAPLE was generated by homology in combination with protein-protein docking. First, a model of the DAPLE GBA motif (residues 1663–1680) was built via homology to GIV residues 1673–1690 using an X-ray co-crystal structure of rat $G\alpha_{i3}$ bound to a human GIV peptide (PDB code 6MHF) as a template (35). The position of hydrogens and the isomeric/tautomeric state and positioning of side chains were energetically optimized on the parent structure prior to modeling. The binding of the GBA motif of DAPLE to the $G\alpha_{i3}$ structure (PDB code 6MHF, chain A) was then simulated with a rigid body two-stage fast Fourier transform protein-protein docking procedure. The solution with the lowest predicted energy was selected, and the sidechain positions of DAPLE residues were energetically minimized with respect to the binding site on $G\alpha_{i3}$. Homology modeling and protein docking were performed with ICM version 3.8 – 3 (Molsoft LLC., San Diego) (84, 85). Model images were

generated with PyMOL Molecular Graphics System, version 2.3.0 (Schrodinger, LLC.).

Peptide synthesis

Peptides were synthesized as described previously (20, 24). The sequences corresponding to the GBA motif of human DAPLE (residues 1662-1695, SASPSSEMVTLEEFLEESNR-SSPTHDTPSCRDDL) or control peptide, corresponding to the GBA motif of GIV 1671–1705 containing the mutation F1685A (KTGSPGSEVVTLQQALEESNKLTSVQIKSSSQENL) were synthesized using the in situ neutralization protocol for t-butoxycarbonyl-solid-phase peptide synthesis on a p-methylbenzhydrylamine resin (Novabiochem, 0.67 mmol/g, 100-200 mesh). Following chain elongation, peptides were cleaved using a solution of hydrofluoric acid containing 5% anisole for 1 h at 0 °C. Next, the hydrofluoric acid solution containing the peptides was removed under vacuum, and the solution was crushed out with Et₂O and filtered. The collected solids were redissolved in a 50% CH₃CN/H₂O solution containing 0.05% of trifluoroacetic acid (TFA), frozen down, and lyophilized. Crude peptides were purified by reverse phase-HPLC using a Jupiter Proteo (90 Å, 10 μ m, 100 \times 21.2 mm) at a flow rate of 16 ml/min using H₂O (A, 0.1% TFA) and CH₃CN (B, 0.05% TFA) as eluents following a linear gradient: from 5% B to 70% B in 80 min. The identity and final purity (>97%) of the peptides were determined by analytical RP-HPLC and MS (ESI-TOF).

Steady-state GTPase assays

Steady-state GTPase assays of $G\alpha_{i3}$ and $G\alpha_{s}$ were performed as described previously (42, 82). Briefly, His-G α_{i3} (100 nm) or His-G α_s (100 nm) was diluted in assay buffer (20 mm Na-HEPES, pH 8, 100 mm NaCl, 1 mm EDTA, 2 mm MgCl₂, 1 mm DTT, 0.05% (w/v) $C_{12}E_{10}$) and preincubated with peptides or purified proteins (as indicated in the figures) for 15 min at 30 °C for $G\alpha_{i3}$ or 30 min at 4 °C for $G\alpha_{s}$. Reactions were initiated at 30 °C by mixing (1:1) (v/v) the mixture prepared above with 1 μ M [γ - 32 P]GTP (\sim 50 cpm/fmol) diluted in assay buffer. Duplicate aliquots (25 μ l) were removed at the times indicated in the figures or figure legends, and the reaction was stopped by the addition of 975 μ l of ice-cold 5% (w/v) activated charcoal in 20 mm H₃PO₄, pH 3. Samples were centrifuged for 10 min at $10,000 \times g$, and 500 μ l of the resultant supernatants were scintillation counted to quantify the amount of [32P]P_i released. Background [32P]P; detected in the absence of G protein was subtracted from each reaction. Data are expressed as picomoles of $[\gamma^{-32}P]P_i$ released or percentage of $[\gamma^{-32}P]P_i$ released compared with control. EC₅₀/IC₅₀ values were estimated from sigmoidal dose-response curve fits generated using the GraphPad software.

GTP_{\gamma}S-binding assays

GTP γ S-binding assays for $G\alpha_{i3}$ and $G\alpha_s$ were performed essentially as described previously (42, 82). Purified His- $G\alpha_{i3}$ (100 nm) and His- $G\alpha_s$ (100 nm) were diluted in assay buffer (20 mm Na-HEPES, pH 8, 100 mm NaCl, 1 mm EDTA, 25 mm MgCl_{2} , 1 mm DTT, 0.05% (w/v) $\mathrm{C}_{12}\mathrm{E}_{10}$) and preincubated with peptides or purified proteins (as indicated in the figure) for 15 min at 30 °C for $G\alpha_{i3}$ or 30 min at 4 °C for $G\alpha_{s}$. Reactions were



initiated by adding an equal volume of assay buffer containing 1 $\mu\rm M$ [$^{35}\rm S$]GTP $\gamma\rm S$ (\sim 50 cpm/fmol) at 30 °C for G α_{i3} or 20 °C for G α_s . Duplicate aliquots (25 $\mu\rm l$) were removed at different time points as described in the figures, and binding of radioactive nucleotide was stopped by addition of 3 ml of ice-cold wash buffer (20 mM Tris-HCl, pH 8.0, 100 mM NaCl, 25 mM MgCl₂). The quenched reactions were rapidly passed through BA-85 nitrocellulose filters (GE Healthcare) and washed with 4 ml of cold wash buffer. Filters were dried and subjected to liquid scintillation counting. Background [$^{35}\rm S$]GTP $\gamma\rm S$ detected in the absence of G protein was subtracted from each reaction. Data are expressed as picomoles of [$^{35}\rm S$]GTP $\gamma\rm S$ bound or percentage of [$^{35}\rm S$]GTP $\gamma\rm S$ binding relative to control (% of control).

GTP γ S-binding assays with $G\alpha_{\alpha}$ were performed using two different buffers: assay buffer A, containing (NH₄)₂SO₄ (20 mm Na-HEPES, pH 7.5, 100 mm NaCl, 0.1 mm EDTA, 0.3 mm $MgCl_2$, 0.2 M (NH₄)₂SO₄, 1 mm DTT, 0.05% (w/v) $C_{12}E_{10}$), and assay buffer B, not containing (NH₄)₂SO₄ (20 mm Na-HEPES, рН 7.5, 100 mм NaCl, 1 mм EDTA, 10 mм MgCl₂, 1 mм DTT, 0.05% (w/v) Genapol). Purified $G\alpha_{\rm q}$ (100 nm) was diluted in either assay buffer A or assay buffer B and preincubated at 4 °C for 30 min with peptides or purified proteins, as indicated in the figure legends. Reactions were initiated by adding an equal volume of assay buffer containing 0.5 μ M [35 S]GTP γ S (\sim 50 cpm/ fmol) at 20 °C. Single aliquots (25 µl) were removed at different time points (as indicated in the figures), and reactions were stopped and analyzed as described above. EC₅₀/IC₅₀ values were estimated from sigmoidal dose-response curve fits using the GraphPad software. The rate constants of time-course experiments were estimated by fitting the data to a one-phase exponential association curve using GraphPad.

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