

Prediction of Protein–Protein Interfaces on G-Protein β Subunits Reveals a Novel Phospholipase C β 2 Binding Domain

Erin J. Friedman¹, Brenda R. S. Temple^{2,3*}, Stephanie N. Hicks⁴,
John Sondek^{3,4,5}, Corbin D. Jones^{1,6} and Alan M. Jones^{1,4}

¹Department of Biology,
University of North Carolina
at Chapel Hill, Chapel Hill,
NC 27599, USA

²The R. L. Juliano Structural
Bioinformatics Core Facility,
University of North Carolina
at Chapel Hill, Chapel Hill,
NC 27599, USA

³Department of Biochemistry
and Biophysics, University
of North Carolina School of
Medicine, Chapel Hill,
NC 27599, USA

⁴Department of Pharmacology,
University of North Carolina
School of Medicine, Chapel Hill,
NC 27599, USA

⁵Lineberger Comprehensive
Cancer Center, University
of North Carolina School
of Medicine, Chapel Hill,
NC 27599, USA

⁶Carolina Center for Genome
Sciences, University of North
Carolina at Chapel Hill,
Chapel Hill, NC 27599, USA

Received 10 February 2009;
received in revised form
8 July 2009;
accepted 27 July 2009
Available online
30 July 2009

Edited by M. Sternberg

G β subunits from heterotrimeric G-proteins (guanine nucleotide-binding proteins) directly bind diverse proteins, including effectors and regulators, to modulate a wide array of signaling cascades. These numerous interactions constrained the evolution of the molecular surface of G β . Although mammals contain five G β genes comprising two classes (G β 1-like and G β 5-like), plants and fungi have a single ortholog, and organisms such as *Caenorhabditis elegans* and *Drosophila melanogaster* contain one copy from each class. A limited number of crystal structures of complexes containing G β subunits and complementary biochemical data highlight specific sites within G β s needed for protein interactions. It is difficult to determine from these interaction sites what, if any, additional regions of the G β molecular surface comprise interaction interfaces essential to G β 's role as a nexus in numerous signaling cascades. We used a comparative evolutionary approach to identify five known and eight previously unknown putative interfaces on the surface of G β . We show that one such novel interface occurs between G β and phospholipase C β 2 (PLC- β 2), a mammalian G β interacting protein. Substitutions of residues within this G β –PLC- β 2 interface reduce the activation of PLC- β 2 by G β 1, confirming that our *de novo* comparative evolutionary approach predicts previously unknown G β –protein interfaces. Similarly, we hypothesize that the seven remaining untested novel regions contribute to putative interfaces for other G β interacting proteins. Finally, this comparative evolutionary approach is suitable for application to any protein involved in a significant number of protein–protein interactions.

© 2009 Elsevier Ltd. All rights reserved.

Keywords: heterotrimeric G-proteins; phospholipase C- β 2; protein surface evolution; interface prediction; PLC- β 2–G β γ interaction interface

*Corresponding author. The R. L. Juliano Structural Bioinformatics Core Facility, University of North Carolina at Chapel Hill, Genetic Medicine Building, 120 Mason Farm Road, Suite 3010, Campus Box 7260, Chapel Hill, NC 27599-7260, USA. E-mail address: btemple@med.unc.edu.

Abbreviations used: G-protein, guanine nucleotide-binding protein; GRK, G-protein-coupled receptor kinase; MSA, multiple sequence alignment; PLC- β 2, phospholipase C β 2; RGS9, regulator of G-protein signaling 9; ROI, region of interest; RSA, relative solvent accessibility.

Introduction

Gene duplication is a fundamental source of genetic and phenotypic novelty.¹ After duplication, one of the new paralogs is freed from functional constraint, enabling it to evolve new functions. For genes encoding proteins that interact with other proteins, this process often liberates one copy to develop a new set of interactions. This is particularly true for signaling molecules that are used by organisms to communicate between cells and to perceive their environment. For example, evolution of increasingly complex organisms correlates with the enormous diversification of heterotrimeric G-protein (guanine nucleotide-binding protein) signaling complexes and cell surface G-protein-coupled receptors. The G-protein complex consists of a heterotrimer composed of G α , G β , and G γ subunits. Upon activation by cell surface receptors, the complex dissociates into a free G α subunit and a G $\beta\gamma$ dimer, both of which bind to and signal through other proteins. Signaling typically terminates when the heterotrimer reforms.^{2,3} *Saccharomyces cerevisiae* has two G α genes but only one each of G β and G γ genes; mammalian genomes encode 16 G α , 5 G β , and 12 G γ genes.² This complex array of subunit combinations allows for diverse signaling possibilities. It was previously thought that G α was the primary signaling molecule in mammals, while the sole function of G β was to inhibit G α signaling and to provide for its membrane localization.⁴ It is now clear that G β also modulates downstream targets; a subset of signaling pathways is uniquely regulated by the G β subunit.⁵

Several proteins that bind G β have been identified in mammals. In addition to G α and G γ , interacting proteins such as the localization chaperone phosducin, G-protein-coupled receptor kinases (GRKs), phospholipase C β 2 (PLC- β 2), regulator of G-protein signaling 9 (RGS9), calcium channels, potassium channels, and adenylyl cyclase 2 bind G β proteins. Mammalian G β interacting proteins arose differentially over evolutionary time as not all eukaryotes contain homologs to all known interactors.² For example, G α and phosducin are present in all eukaryotes beginning with plants. Canonical RGS9, PLC- β 2, and GRK2 originated and were maintained in metazoans at least by the time of the formation of annelids since *Caenorhabditis elegans* contains bona fide RGS9, PLC- β 2, and GRK2 orthologs.^{6–8} As the G β subunit acquired additional binding partners throughout evolution, new G β -protein interfaces likely evolved to accommodate these interactions. These new interfaces may have partially overlapped with existing interfaces since, for example, the interface from many G β interactors overlaps with the G α -G β interface.⁹ However, these interfaces may have also utilized regions on the G β molecular surface that previously had no associated function.

Two major experimental approaches, structural and biochemical studies, have characterized some binding interfaces between G β and interacting

proteins. Currently, there are four crystal structures of mammalian G β subunits in complex with signaling proteins: G β 1 γ 1-G α ,¹⁰ G β 1 γ 2-GRK2,¹¹ G β 1 γ 1-phosducin,¹² and G β 5-RGS9.¹³ These structures provide a three-dimensional view of where proteins interact but provide limited information regarding the importance of individual contacts. Additionally, structures of G β in complex with peptides^{14,15} provide partial information on physiologically relevant interfaces within G β . Biochemical studies include both targeted mutational studies^{9,16,17} and targeted domain-swapping or peptide-binding experiments.^{18–20} Mutational studies directed by the structural studies are limited by the number of available structures. Interpretation of these studies is complicated by overlap between the implicated binding regions. When crystal structures of a particular G β complex are not available, mutations are frequently targeted to an interface identified in a solved G β complex structure. For example, Ford *et al.*⁹ created several mutations in the G α binding interface to show that this interface was utilized in part by five other interacting proteins. This type of approach however does not elucidate binding sites, or even those regions of the binding interface, that are not shared with the G α interface. To locate these alternative sites, groups such as Panchenko *et al.*¹⁷ performed mutational analyses outside of known binding areas. Although this study identified mutant regions of G β with reduced ability to activate PLC- β 2, much refinement remains necessary to identify individual critical interaction sites. Moreover, this study left many regions of the G β surface unexplored.

Bioinformatic analyses have been developed to identify functional residues, including those comprising conserved patches on the surface of a protein that may function as a binding interface. Three such analyses include evolutionary trace,²¹ DIVERGE,^{22,23} and ConSurf.²⁴ All three rely on multiple sequence alignments (MSAs) and phylogenetic trees to identify structurally clustered functional residues. Evolutionary trace has been applied to G β and the two predicted interfaces correlated with those of G α and G γ .²⁵ To date, none of these three methods has predicted previously uncharacterized G β interfaces.

We took advantage of both the rich source of divergent G β subunit sequences and the differential evolutionary emergences of known G β interactors to understand not only how to predict novel interaction interfaces on the G β surface in the absence of crystallized complexes but also how these new interactions gave rise to new binding surfaces. These results can be utilized for more targeted and informed biochemical studies. Based on the hypothesis that the acquisition of mammalian-like sequence identity on the surface of G β reflects the utilization of new binding interfaces, we applied a suite of bioinformatic and phylogenetic techniques to follow the shift in patterns of amino acid conservation in G β s, concentrating on changes between distinct points in the evolutionary history of the G β subunit represented by five reference

species. By placing this conservation in a structural context, we predicted regions of interest (ROIs) that are composed of adjacent surface residues that simultaneously evolved to residues conserved with mammals. We identified a novel PLC- β 2 interface by demonstrating that at least one ROI that became conserved in *C. elegans* is involved in PLC- β 2 activation by G β . Similarly, we propose that the remaining ROIs also compose at least portions of binding interfaces.

Results and Discussion

G β proteins fall into two major classes: G β 1-like and G β 5-like

Since most extant plant and fungal species have a single G β , while nematodes and later metazoans have at least two G β subunits, the first G β gene duplication occurred between the splitting of fungi and *C. elegans* from the mammalian lineage. To compare preduplication plant and fungal G β sequences with extant postduplication G β sequences from metazoans, we reconstituted ancestral sequences to plants and to fungi from MSAs (Figs. S1 and S2, respectively) of extant plant and fungal G β sequences using the Bayesian ancestral reconstruction implemented in MrBayes (see [Materials and Methods](#)). These inferred preduplication ancestors were important as they, and not individual plant or fungal species, reflect sequence constraints common to all plants or fungi and therefore more closely reflect the predecessor of all extant postduplication G β genes.

An MSA containing G β sequences from the plant ancestor, fungal ancestor, and extant *C. elegans*, *Drosophila melanogaster*, and human G β sequences was created (Fig. S3) for comparison of preduplication and postduplication G β genes. This MSA was used to generate a Bayesian phylogenetic tree (Fig. S4) that elucidates the evolutionary relationship between the different G β proteins. Human G β 1, G β 2, G β 3, and G β 4 sequences formed a monophyletic G β 1-like clade also containing *D. melanogaster* and *C. elegans* G β 1 sequences. Human G β 5 formed a G β 5-like monophyletic clade containing *D. melanogaster* G β 5 and *C. elegans* G β 2 sequences. Both plant and fungal ancestors were outside of these two clades.

Based on the phylogenetic data (Fig. S4) and corroborating previous observations that G β 5 often behaves differently from the other four mammalian G β proteins,⁵ metazoan G β proteins can be grouped into two major classes, G β 1-like and G β 5-like. The initial G β gene duplication between fungi and *C. elegans* gave rise to the G β 5-like family since *C. elegans* is the earliest species examined to contain a G β 5-like protein. The ancestral plant and fungal G β proteins each contains characteristics of both major classes, but neither strictly belongs to either class. Finally, the phylogeny revealed that, after duplication, the G β 5-like genes diverged from the ancestor

more than the G β 1-like genes, while the G β 1-like genes maintained more ancestral characteristics.

Interaction interfaces identified from G β complex structures co-evolved with interactors

Complexes containing mammalian G β 1-G α , G β 1-GRK2, G β 1-phosducin, and G β 5-RGS9 are shown in Fig. 1 (G β in spheres and interacting protein in ribbons). While these structures provide a three-dimensional view of G β -protein complexes, the interfaces between the proteins must still be defined.²⁶ To define sites of interaction, we calculated the solvent accessibility of each residue in the four crystal structures using Naccess.²⁷ Those G β residues whose side chains have lower relative solvent accessibility (RSA) by more than 5% between the monomer and the protein complex, including those that hydrogen bond with the binding partner, were defined to comprise the interface and were noted on the structure (see [Materials and Methods](#)) (Fig. 1). These four G β interacting proteins (1) cover three major areas on the G β surface, (2) all partially overlap with one another, and (3) appeared at different times over G β evolution.

In order to examine the evolution of these known G β -protein interfaces, we compared the patterns of amino acid conservation within these regions. For each interface residue in the MSA, comparisons were made between each of the four reference sequences and the corresponding mammalian residue. For G β 1 complexes (G α , GRK2, and phosducin), the plant ancestor, fungal ancestor, *C. elegans* G β 1, and *D. melanogaster* G β 1 were compared with bovine G β 1. Identical interface residues were mapped onto the bovine G β 1 structure (1got.pdb). For the RGS9 complex, the plant ancestor, fungal ancestor, *C. elegans* G β 5, and *D. melanogaster* G β 5 were compared with mouse G β 5 and identities were noted on the mouse G β 5 structure (2pbi.pdb). As expected, conservation within known binding areas correlated with the known utilization of these regions (Fig. 2). The complexes subsequently discussed were observed in the four crystal structure complexes.

G β 1-G α

The G β -G α interaction is the most ancient; thus, it would be expected that the G α -G β interface is well conserved in all organisms studied. As expected, the G α binding interface was highly conserved (77%–100% identity) in all sequences analyzed (Fig. 2a, top row) as illustrated by the predominantly orange coloration of the interface.

G β 1-GRK2

The GRK2 interface (Fig. 2a, second row) was completely conserved in *C. elegans* and *D. melanogaster*, correlating with the emergence of GRK2 between fungi and *C. elegans*. Although a large portion of the interface was also conserved in the

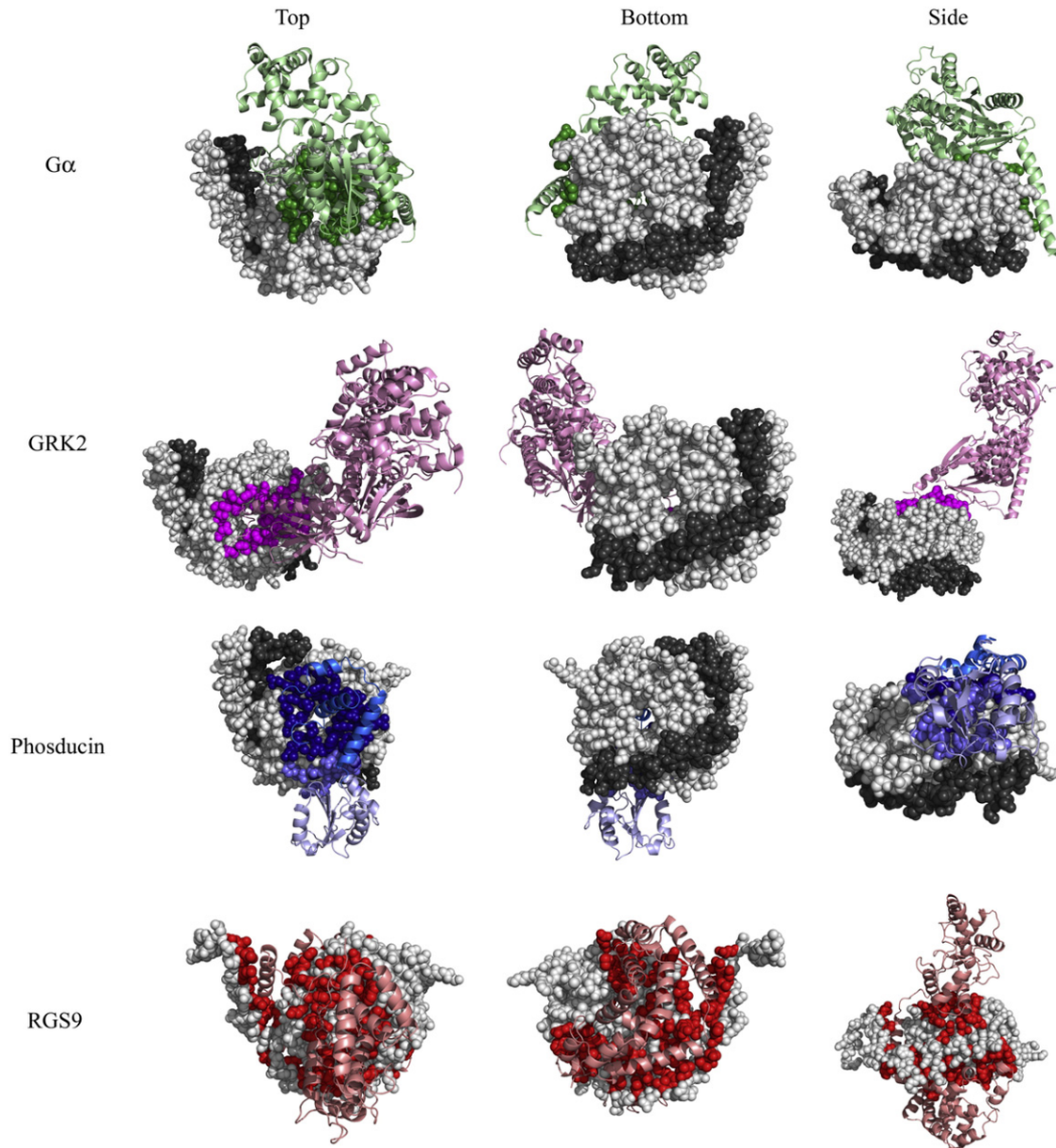


Fig. 1. Binding interfaces of four G β interacting proteins as determined by crystal structures. Top, bottom, and side views of G β or the G β γ dimer (G β in light gray spheres and G γ in dark gray spheres) bound to four interacting proteins (ribbon): G β 1–G α in green (1got.pdb), G β 1–GRK2 in magenta (1omw.pdb), G β 1–phosducin in blue (N-terminal domain is dark and C-terminal domain is light; 1a0r.pdb), and G β 5–RGS9 in red (2pbi.pdb). G β residues that contact each interacting protein are colored accordingly. Binding contacts were determined by evaluating the solvent accessibility difference between single molecules and those in complex.

plant (62%) and fungal (92%) ancestors, this can be attributed to the fact that the interface largely overlaps that of G α .

G β 1–phosducin

Plants contain several phosducin-like sequences,²⁸ and our yeast three-hybrid data suggest that at least one phosducin in *Arabidopsis thaliana* interacts with the *A. thaliana* G β (Fig. 3). The phosducin interface is highly conserved in *C. elegans* and *D. melanogaster* (94% identity); however, it is only partly conserved in plant and fungal ancestors (55% and 64%,

respectively). Mammalian phosducin has two domains: a helical N-terminal domain and a C-terminal domain (see Fig. 1 for locations of N- and C-terminal binding interfaces), both of which bind mammalian G β .²⁹ The plant and fungal ancestors show high conservation in the N-terminal binding area (68% and 76%, respectively) but not in the C-terminal binding area (25% and 38%, respectively) (Fig. 2a, third and fourth rows). Two possible explanations for these results are that plants and fungi bind only one phosducin domain and not the second and that the second domain binds in a species-specific manner.

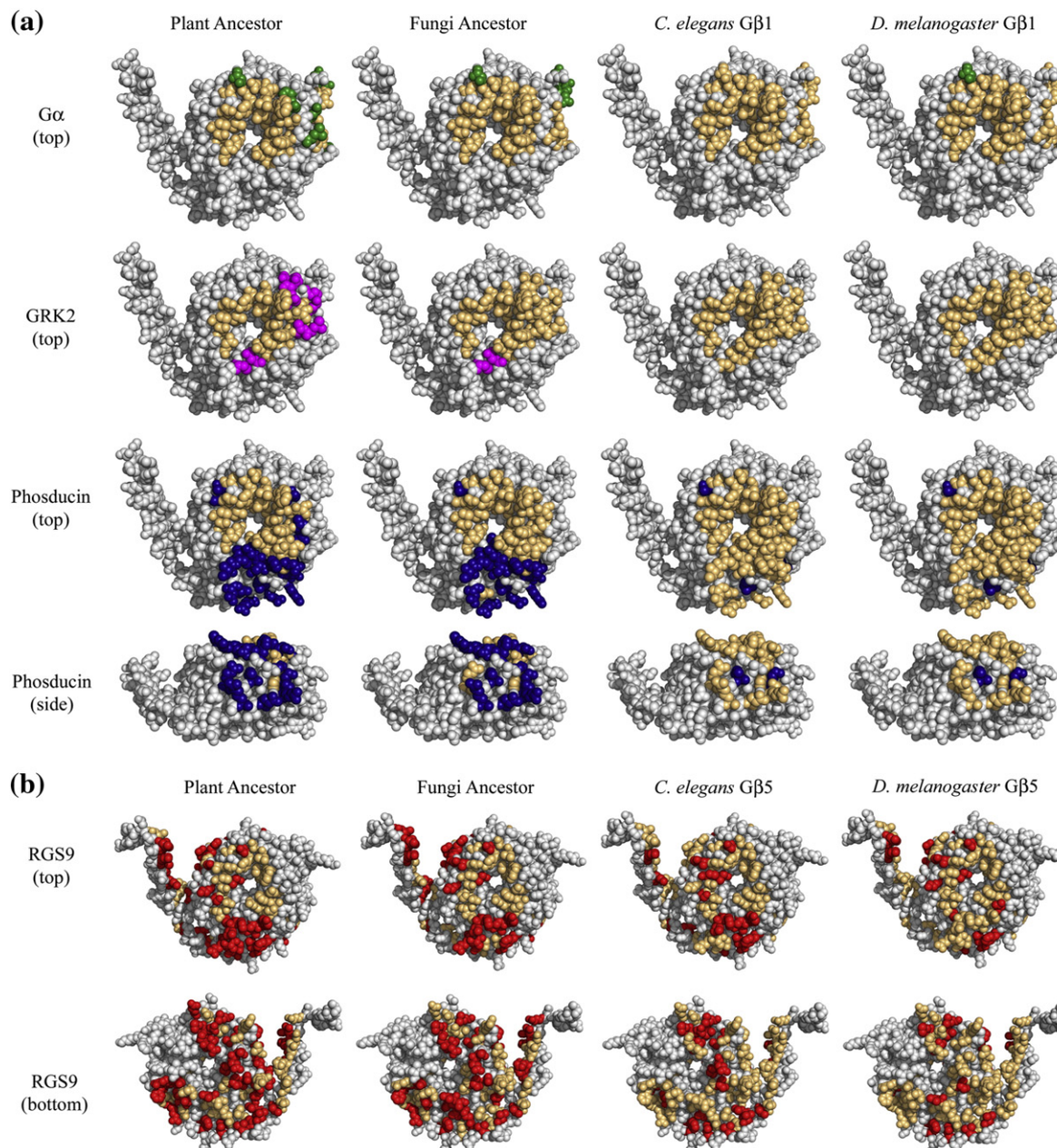


Fig. 2. Conservation within known binding interfaces based on bovine G β 1 crystal structures. (a and b) Conservation of the (a) G α , GRK2, and phosducin binding interfaces on G β 1 (1got.pdb) and (b) RGS9 binding interface on G β 5 (2pbi.pdb) in four reference organisms. While the interface is composed of all colored residues, conserved residues are colored light orange and nonconserved residues are colored green (G α), blue (phosducin), magenta (GRK2), or red (RGS9). Only residues in the binding area were analyzed.

G β 5–RGS9

The RGS9 binding interface (Fig. 2b) is only 42% identical in the plant ancestor and 50% identical in the fungal ancestor. Correlating with the genesis of the RGS9 protein, conservation in the binding area rises to 67% and 76% in *C. elegans* and *D. melanogaster*, respectively. Although this is markedly less than the 90%–100% identity seen in corresponding G β 1 binding areas, it is important to note the size of the RGS9 interface. Encompassing 103 residues, the RGS9 interface is over three times larger than the G α (31 residues) and GRK2 (26 residues) interfaces and twice

the size of the phosducin interface (47 residues). Thus, we expect that the RGS9 interface could tolerate more substitutions than its smaller counterparts and that only those residues making energetically critical contacts in the interaction interface are conserved.

Newly conserved regions of adjacent surface residues arose over time and likely contribute to G β –protein interfaces

As conservation within known G β –protein interfaces correlated with the emergence of proteins

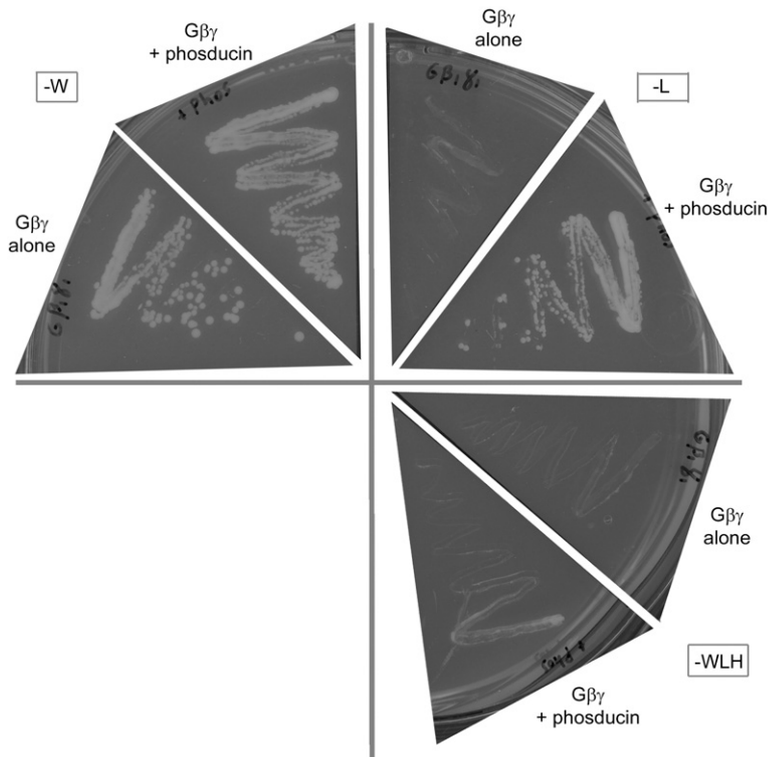


Fig. 3. An *Arabidopsis* phosducin and G β interact physically. Growth of yeast strain AH109 containing the genes indicated (AtG β 1 γ 1 alone or AtG β 1 γ 1 and phosducin) on yeast dropout medium. Medium missing tryptophan (–W) selects for the AtG β 1 γ 1 vector, resulting in positive growth for both genotypes. Medium missing leucine (–L) selects for the phosducin vector, resulting in no growth for the strain lacking phosducin and positive growth for the strain containing phosducin. Medium missing tryptophan, leucine, and histidine (–WLH) selects for a positive interaction between the two genes, resulting in no growth for the strain containing AtG β 1 γ 1 alone and positive growth for the strain containing both AtG β 1 γ 1 and phosducin. The latter growth indicates that the two genes physically interact.

utilizing these interfaces, we next used a similar method of analysis to predict *novel* binding interfaces. This unbiased comparative evolutionary analysis was not restricted to predetermined binding interfaces and was performed on the entire molecule, but it was otherwise similar to the methods used in the validation stage described in the previous section. Bovine G β 1 was compared at each residue in the MSA with the plant ancestor, fungal ancestor, *C. elegans* G β 1, and *D. melanogaster* G β 1 (Fig. 4a). Identical residues between the plant ancestor and bovine G β 1 were located on the surface of G β 1 and colored dark green. These 162 residues form region zero (ROI0) and represent the primordial function of the G β molecule, since the plant molecule is the most ancestral-like of the G β family. For all other reference species, conserved residues that continued to be conserved from the previous species were colored light green (indicating the persistence of an existing function), while newly conserved residues were colored dark green (indicating the emergence of potential novel function). Similarly, mouse G β 5 was compared with the plant ancestor, fungal ancestor, *C. elegans* G β 5, and *D. melanogaster* G β 5 and conserved residues were mapped onto the structure (Fig. 4b).

Due to its primordial nature, ROI0 contains residues conserved both for structural maintenance and for interaction surfaces. ROI0 contains the residues (e.g., W99, M101 K57, Y59, L117, D186, D228, and W332) previously indicated as interacting with multiple effectors and that also made energetically critical contacts in various interaction interfaces.^{9,14} Subsequent non-ancestral ROIs in each organism (for both G β 1 and G β 5) were

defined as clusters of three or more structurally adjacent (within 5 Å³⁰) newly conserved surface residues and are encircled on the structure (Fig. 4; for residue designations, see Table 1). It is important to note that although all surface-exposed residues could form part of an interface, we chose a larger cluster size (containing three or more newly fixed residues) in order to predict clusters that likely make significant energetic contributions to binding. We also ignored several large clusters near the G γ binding area due to their likely involvement in the G β –G γ interface.

Five of the 13 ROIs (ROI7, ROI10, ROI11, ROI12, and ROI13) could be explained by existing structural data. ROI7 lies in the phosducin binding region on G β 1 (Fig. 1). Portions of ROI10–ROI13 localize to the RGS9 binding interface on G β 5 (Fig. 1). Additionally, a portion of ROI2 was shown to interact with G α , adenylyl cyclase 2, calcium channels, potassium channels, and PLC- β 2 in mutational studies.⁹ These data support the hypothesis that regions of adjacent surface residues that become conserved to the mammalian state at the same time are likely sites of G β –protein interactions. Similarly, we hypothesize that the remaining ROIs are binding interfaces with yet-to-be identified interacting proteins or additional, undescribed binding interfaces with previously identified interacting proteins.

Several ROIs contain G β residues that activate PLC- β 2

To test our hypothesis that ROIs represent G β –protein interaction interfaces, we mutated residues

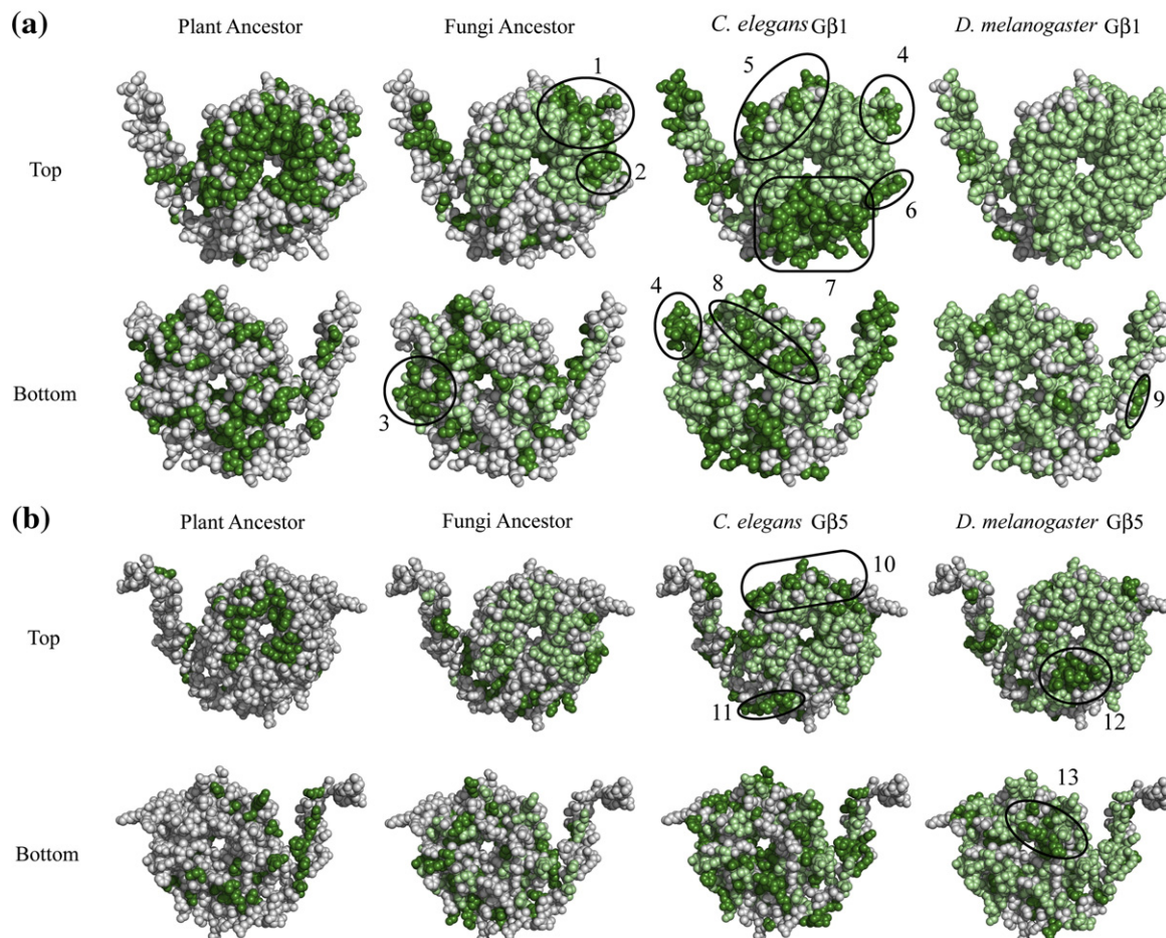


Fig. 4. G β ROIs as determined by a comparative evolutionary analysis. Three views (top, bottom, and side) of the evolution of conserved regions in the plant ancestor, fungal ancestor, *C. elegans* G β , and *D. melanogaster* G β [(a) G β 1 (1got.pdb) and (b) G β 5 (2pbi.pdb)]. The dark green regions of the plant ancestor highlight primordial function and form ROI0. In all other organisms, newly conserved residues (those that matched the mammalian value) are colored dark green, while conserved residues present in a previous organism are colored light green. Thus, dark green patches represent acquisition of new function. ROIs are indicated and corresponding residues are listed in [Table 1](#).

likely to interact with PLC- β 2. Despite the absence of a crystal structure of the G $\beta\gamma$ -PLC- β 2 complex, several studies implicated multiple potential interfaces. Peptide-binding assays implicated a region at the base of G β 's N-terminal helix.^{18,20} A patch of residues near this region was implicated as an interface when it was revealed that the C-terminal tail of G γ was responsible for the activation of PLC- β 2.¹⁹ Therefore, ROI4, ROI6, and ROI7 were chosen for mutational studies due to their proximity to the general region implicated in these peptide-binding and domain-swapping experiments. Additionally, all three regions became conserved between the fungi and *C. elegans* split, corresponding with the emergence of PLC- β proteins.

In order to evaluate the extent of PLC- β 2 activation, we transfected COS-7 cells with wild-type and mutant forms of G β 1 and wild-type G γ 2 in the presence or absence of PLC- β 2. Equal expression of all G β 1, G γ 2, and PLC- β 2 constructs was confirmed by immunoblot analysis. Additionally, equivalent expression of G γ 2 was observed in the presence of all forms of G β 1. The location of

each mutation on the surface of G β 1 is depicted in [Fig. 5](#). The construct containing the double mutation T65A_D322A, which is located outside of the predicted binding region and lacks sequence conservation with bovine G β 1, activates PLC- β 2 to the same extent as wild-type G $\beta\gamma$. Conversely, two single mutations, W99A and D228A, abolish PLC- β 2 activation as previously shown.⁹ Within the newly predicted binding areas, the mutations that most significantly reduced PLC- β 2 activation were found in ROI4 and ROI6 ([Figs. 4a](#) and [5](#)). R52S_F335S slightly reduced PLC- β 2 activation, while K127S_R129S_E130S nearly abolished PLC- β 2 activation. In summary, when residues within the new binding areas predicted by our *de novo* analysis are mutated, there is a concordant drop in activity; conversely, when residues we identified as not part of binding surfaces are mutated, there is no change in activation. These results indicate that our evolutionary structural analysis predicts the nonoverlapping portions of interaction interfaces and can be used to further dissect the PLC- β 2 interface on G β 1.

Table 1. ROI residue numbers

ROI	Residues	First conserved in
1	R96, S97, S98, I120, R134, E138, E172	Fungi Gβ1
2	L55, A56, S334	Fungi Gβ1
3	D66, R68, Y85, N88, V90, Y105	Fungi Gβ1
4	K127, R129, E130, N132	<i>C. elegans</i> Gβ1
5	T143, T173, Q175, Q176, T181, T184, M217	<i>C. elegans</i> Gβ1
6	R46, T47, R52, D312, F335	<i>C. elegans</i> Gβ1
7	R42, Q44, D267, N268, I269, I270, C271, G272, I273, D290, D291, N293, N295, V307, A309	<i>C. elegans</i> Gβ1
8	L152, D153, N155, D195, R197, L210	<i>C. elegans</i> Gβ1
9	D20, A24	<i>D. melanogaster</i> Gβ1
10	T102, P104, T106, N141, K146, N154	<i>C. elegans</i> Gβ5
11	E43, K279, E280, S281	<i>C. elegans</i> Gβ5
12	F284, N303, Y305	<i>D. melanogaster</i> Gβ5
13	N163, L203, P205, E207	<i>D. melanogaster</i> Gβ5

Shown are mammalian residue positions for each Gβ residue within the ROIs highlighted in Fig. 4.

Molecular evolution can be used as a tool to predict novel binding interfaces

Conservation is a hallmark of a residue's functional importance,²³ and as the plant ancestor is most similar to the most ancient Gβ molecule, primordial Gβ function can be inferred to correlate with those residues that were conserved in the plant ancestor (ROI0) (Fig. 4). These primordial functional regions comprise both conserved protein-binding interfaces and regions of the protein important for structural integrity. With each comparative step (e.g., between plants and fungi, fungi and *C. elegans*,

etc.), new functional regions became conserved. These regions may represent sites of novel Gβ–protein interaction or new interaction sites with existing proteins (co-evolution). Lineage-specific protein interactions will however not be revealed by this type of analysis. Lineage-specific interactions would be indicated by conservation of a residue within a particular organismal group (e.g., within all plants or within all fungi) that is *not* conserved outside of that group (e.g., between plants and fungi). All interfaces identified in our analysis are ones that are maintained in the mammalian lineage. Although species-specific interactions are also of great interest, a different approach would be necessary to reveal those regions.

The Gβ gene family was produced by a series of gene duplications. After these events, the new Gβ paralogs likely evolved novel functions as a result of relaxed functional constraint. Eventually, as these new Gβ functions become critical to the organism, these new functional regions become highly conserved regions of the Gβ structure and include the ROIs we identified. We hypothesize that these ROIs represent regions of newly acquired function in the organism in which they first appeared. For instance, portions of six ROIs presented in Fig. 4 (ROI2, ROI7, and ROI10–ROI13) contain residues that likely bind interactors as previously identified in mutational or structural studies. The remaining ROIs in Fig. 4 are in regions that lie outside of known regions presented thus far. Although interactions with many proteins have been determined by mutational studies, many studies focus on known interfaces and therefore indirectly introduce bias. For example,

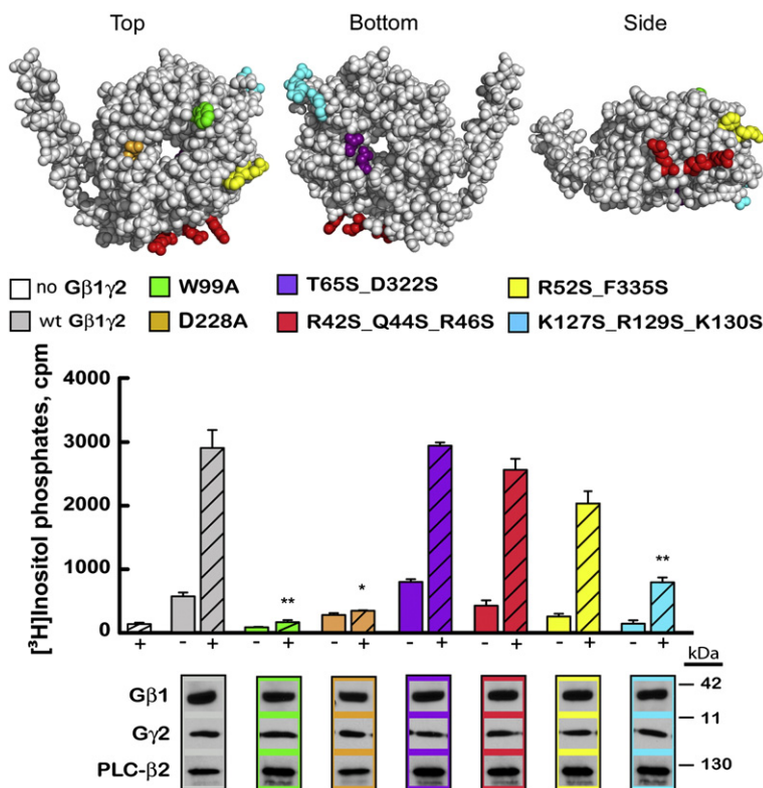


Fig. 5. Regions of Gβ involved in PLC-β2 activation. Groups of residues mutated on the Gβ surface (1got.pdb) are colored and coordinate with colors in the bar graph and associated blots. Wild-type and mutant Gβ1 subunits were tested in the absence (-) or presence (+) of PLC-β2 for their ability to activate PLC-β2 (**p*<0.01 and ***p*<0.005; error bars represent standard error). Immunoblot analysis confirmed equal expression of all proteins utilized.

since the G α binding area is utilized in many G β -protein interactions, studies historically focused on these residues. Our *de novo* analysis eliminates this bias, generating a set of potential binding interfaces that are yet to be explored by mutational studies. As with the residues implicated in the PLC- β 2 binding interface, these ROIs would make ideal targets for site-directed mutagenesis. In this manner, we can identify new G β interacting proteins and further characterize interactions with existing proteins. Ultimately, this will allow us to characterize new G β -mediated signaling pathways.

Materials and Methods

Ancestral reconstruction

To reconstruct plant and fungal ancestors, we queried sequence databases with known plant and mammalian G β sequences for full-length plant and fungal G β homologs. Additionally, G β sequences were also retrieved for two outgroups: Entamoeba (*Entamoeba dispar*) and diatom (*Thalassiosira pseudonana*). MSAs were generated in ClustalX.³¹ ExPasy and Genbank IDs for each gene used in the alignment are denoted in Figs. S1 and S2. For each MSA generated, all gaps except those found exclusively in the two outgroups were removed. From the resulting NEXUS file, ancestors were generated with MrBayes^{32,33} using the fixed equalin model, using the inverse gamma rate, and sampling 100,000 generations at a frequency of 100. In order to eliminate ancestral residue values chosen with low confidence, we accepted and included in the ancestral sequence only those residues predicted >90% of the time with a maximum value >0.8. All other residues were assigned a value of "X" as they were too variable to be called with confidence.

Sequence collection, alignment, and phylogeny generation

All G β sequences from *D. melanogaster*, *C. elegans*, and human (ExPasy and Genbank sequence IDs are denoted in Fig. S3) were combined with the plant and fungal ancestral sequences and the diatom and Entamoeba outgroups to create an MSA (see previous section for alignment and NEXUS file generation). The alignments were made robust by structural comparisons between bovine G β 1 and mouse G β 5. MrBayes^{32,33} was run using a fixed equalin model, using the inverse gamma rate, and sampling 1 million generations at a frequency of 100 for three independent runs with a burn-in of 250,000 generations to generate a consensus phylogenetic tree.

Interface determination

In order to identify binding interfaces from structures of G β in complex with interacting proteins, we used Naccess²⁷ to calculate the RSA of all G β residue side chains both as a monomer and in complex with other proteins. All residue side chains with an RSA decrease >5% (RSA of G β monomer-RSA of complex, >5) between monomer and complex were denoted as being present in the interface. This group of residues also included all G β residues that formed hydrogen bonds

with the interacting protein as determined in the PyMOL molecular modeling system.³⁴ The 5% Δ RSA value was chosen because it is more stringent than the commonly utilized 1- \AA^2 change in absolute solvent accessibility^{26,35} but still included all residues that form hydrogen bonds. Additionally, a residue with $\geq 5\%$ RSA is often designated as a surface residue, while a residue with <5% RSA is designated as an interior residue.^{26,35}

Yeast three-hybrid protein interaction

A. thaliana G β (AGB1) and G γ 1 (AGG1) were cloned into the pBridge vector (Clontech, Palo Alto, CA). *A. thaliana* phosducin (At5g14240) was cloned into the p-ENTR/D-TOPO vector (Invitrogen, Carlsbad, CA) and then recombined into the pACTGW-attR Gateway vector,³⁶ which contains an activation domain and is compatible with the pBridge vector. The prey (phosducin) was transformed into yeast strain AH109, which had previously been transformed with the bait (AGB1/AGG1). Both strains (containing the bait alone and containing both bait and prey) were grown on nutritionally selective media. Presence of the bait and that of the prey were confirmed by the expression of nutritional markers (positive growth on media lacking tryptophan and leucine, respectively). Interaction was confirmed by the expression of an additional nutritional marker (positive growth on medium lacking histidine).

Transfection of COS-7 cells with G β 1, G γ 2, and PLC- β 2

The accumulation of [³H]inositol phosphates was measured in transiently transfected COS-7 cells as previously described.³⁷ Briefly, COS-7 cells were plated in 12-well dishes at a cell density of 60,000 cells per well in Dulbecco's modified Eagle's medium supplemented with 10% fetal bovine serum, 10 U/ml of penicillin, and 10 U/ml of streptomycin. Following incubation for 24 h at 37 °C in an atmosphere of 95% air/5% CO₂, cells were transfected with 200 or 300 ng each of the indicated human G β 1 and G γ 2 DNA in the presence and in the absence of 30 ng of PLC- β 2 and empty vector, for a total of 700 ng of DNA per well. DNA was complexed with FuGENE 6 (Roche Applied Sciences, Indianapolis, IN) as per the manufacturer's protocol prior to transfection. Twenty-four hours posttransfection, the medium was aspirated and cells were metabolically labeled with 1 μ Ci/well of *myo*-[2-³H(N)]inositol (American Radiolabeled Chemicals, St. Louis, MO) in inositol-free Dulbecco's modified Eagle's medium for 12–16 h. Subsequently, 10 mM LiCl was added. One hour after incubation with LiCl, reactions were stopped by aspiration of the medium and addition of 50 mM formic acid. Samples were neutralized by the addition of 150 mM NH₄OH, and [³H]inositol phosphates were quantified by Dowex chromatography. Statistical significance for comparisons between the activation of PLC- β 2 by wild-type G β 1 γ 2 and that by each mutant G β γ was determined by performing Student's *t* test assuming equal variance between the log-transformed number of [³H]inositol phosphates for each construct. Western blotting was performed to confirm equal expression of each construct in COS-7 cells using an antibody directed toward the c-Myc epitope (Invitrogen) on human G β 1, the hemagglutinin epitope (Roche Applied Sciences) on G γ 2, and a monoclonal antibody (Santa Cruz Laboratories, Santa Cruz, CA) against PLC- β 2.

Acknowledgements

Work in A.M.J.'s laboratory on the *Arabidopsis* G-proteins was supported by the National Institute of General Medical Sciences (GM065989-01), the U.S. Department of Energy (DE-FG02-05ER15671), and the National Science Foundation (MCB-0718202 and MCB-0723515). Work in J.S.'s laboratory was supported by the National Institutes of Health (5-50921-2311).

Supplementary Data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.jmb.2009.07.076](https://doi.org/10.1016/j.jmb.2009.07.076)

References

- Ohno, S. (1970). *Evolution by Gene Duplication*. Springer, Berlin, Germany.
- Jones, A. M. & Assmann, S. M. (2004). Plants: the latest model system for G-protein research. *EMBO Rep.* **5**, 572–578.
- McCudden, C. R., Hains, M. D., Kimple, R. J., Siderovski, D. P. & Willard, F. S. (2005). G-protein signaling: back to the future. *Cell. Mol. Life Sci.* **62**, 551–577.
- Milligan, G. & Kostenis, E. (2006). Heterotrimeric G-proteins: a short history. *Br. J. Pharmacol.* **147**(Suppl. 1), S46–S55.
- Cabrera-Vera, T. M., Vanhauwe, J., Thomas, T. O., Medkova, M., Preininger, A., Mazzoni, M. R. & Hamm, H. E. (2003). Insights into G protein structure, function, and regulation. *Endocr. Rev.* **24**, 765–781.
- van der Linden, A. M., Simmer, F., Cuppen, E. & Plasterk, R. H. (2001). The G-protein beta-subunit GPB-2 in *Caenorhabditis elegans* regulates the G α -G β signaling network through interactions with the regulator of G-protein signaling proteins EGL-10 and EAT-16. *Genetics*, **158**, 221–235.
- Miller, K. G., Emerson, M. D. & Rand, J. B. (1999). G α and diacylglycerol kinase negatively regulate the G β pathway in *C. elegans*. *Neuron*, **24**, 323–333.
- Fukuto, H. S., Ferkey, D. M., Apicella, A. J., Lans, H., Sharmeen, T., Chen, W. *et al.* (2004). G protein-coupled receptor kinase function is essential for chemosensation in *C. elegans*. *Neuron*, **42**, 581–593.
- Ford, C. E., Skiba, N. P., Bae, H., Daaka, Y., Reuveny, E., Shekter, L. R. *et al.* (1998). Molecular basis for interactions of G protein $\beta\gamma$ subunits with effectors. *Science*, **280**, 1271–1274.
- Lambright, D. G., Sondek, J., Bohm, A., Skiba, N. P., Hamm, H. E. & Sigler, P. B. (1996). The 2.0 Å crystal structure of a heterotrimeric G protein. *Nature*, **379**, 311–319.
- Lodowski, D. T., Pitcher, J. A., Capel, W. D., Lefkowitz, R. J. & Tesmer, J. J. (2003). Keeping G proteins at bay: a complex between G protein-coupled receptor kinase 2 and G $\beta\gamma$. *Science*, **300**, 1256–1262.
- Loew, A., Ho, Y. K., Blundell, T. & Bax, B. (1998). Phosducin induces a structural change in transducin $\beta\gamma$. *Structure*, **6**, 1007–1019.
- Cheever, M. L., Snyder, J. T., Gershburg, S., Siderovski, D. P., Harden, T. K. & Sondek, J. (2008). Crystal structure of the multifunctional G $\beta 5$ -RGS9 complex. *Nat. Struct. Mol. Biol.* **15**, 155–162.
- Davis, T. L., Bonacci, T. M., Sprang, S. R. & Smrcka, A. V. (2005). Structural and molecular characterization of a preferred protein interaction surface on G protein $\beta\gamma$ subunits. *Biochemistry*, **44**, 10593–10604.
- Johnston, C. A., Kimple, A. J., Giguere, P. M. & Siderovski, D. P. (2008). Structure of the parathyroid hormone receptor C terminus bound to the G-protein dimer G $\beta 1\gamma 2$. *Structure*, **16**, 1086–1094.
- Li, Y., Sternweis, P. M., Charnecki, S., Smith, T. F., Gilman, A. G., Neer, E. J. & Kozasa, T. (1998). Sites for G α binding on the G protein β subunit overlap with sites for regulation of phospholipase C β and adenylyl cyclase. *J. Biol. Chem.* **273**, 16265–16272.
- Panchenko, M. P., Saxena, K., Li, Y., Charnecki, S., Sternweis, P. M., Smith, T. F. *et al.* (1998). Sites important for PLC $\beta 2$ activation by the G protein $\beta\gamma$ subunit map to the sides of the β propeller structure. *J. Biol. Chem.* **273**, 28298–28304.
- Bonacci, T. M., Ghosh, M., Malik, S. & Smrcka, A. V. (2005). Regulatory interactions between the amino terminus of G-protein $\beta\gamma$ subunits and the catalytic domain of phospholipase C $\beta 2$. *J. Biol. Chem.* **280**, 10174–10181.
- Myung, C. S., Lim, W. K., DeFilippo, J. M., Yasuda, H., Neubig, R. R. & Garrison, J. C. (2006). Regions in the G protein γ subunit important for interaction with receptors and effectors. *Mol. Pharmacol.* **69**, 877–887.
- Yoshikawa, D. M., Bresciano, K., Hatwar, M. & Smrcka, A. V. (2001). Characterization of a phospholipase C $\beta 2$ -binding site near the amino-terminal coiled-coil of G protein $\beta\gamma$ subunits. *J. Biol. Chem.* **276**, 11246–11251.
- Lichtarge, O., Bourne, H. R. & Cohen, F. E. (1996). An evolutionary trace method defines binding surfaces common to protein families. *J. Mol. Biol.* **257**, 342–358.
- Gu, X. & Vander Velden, K. (2002). DIVERGE: phylogeny-based analysis for functional-structural divergence of a protein family. *Bioinformatics*, **18**, 500–501.
- Zheng, Y., Xu, D. & Gu, X. (2007). Functional divergence after gene duplication and sequence-structure relationship: a case study of G-protein alpha subunits. *J. Exp. Zool. B: Mol. Dev. Evol.* **308**, 85–96.
- Armon, A., Graur, D. & Ben-Tal, N. (2001). ConSurf: an algorithmic tool for the identification of functional regions in proteins by surface mapping of phylogenetic information. *J. Mol. Biol.* **307**, 447–463.
- Lichtarge, O., Bourne, H. R. & Cohen, F. E. (1996). Evolutionarily conserved G $\alpha\beta\gamma$ binding surfaces support a model of the G protein-receptor complex. *Proc. Natl Acad. Sci. USA*, **93**, 7507–7511.
- Jones, S. & Thornton, J. M. (1996). Principles of protein-protein interactions. *Proc. Natl. Acad. Sci. USA*, **93**, 13–20.
- Hubbard, S. J. & Thornton, J. M. (1993). 'NACCESS', computer program. Department of Biochemistry and Molecular Biology, University College London.
- Blaauw, M., Knol, J. C., Kortholt, A., Roelofs, J., Ruchira, Postma, M. *et al.* (2003). Phosducin-like proteins in *Dictyostelium discoideum*: implications for the phosducin family of proteins. *EMBO J.* **22**, 5047–5057.
- Gaudet, R., Bohm, A. & Sigler, P. B. (1996). Crystal structure at 2.4 angstroms resolution of the complex of transducin $\beta\gamma$ and its regulator, phosducin. *Cell*, **87**, 577–588.
- Madabushi, S., Yao, H., Marsh, M., Kristensen, D. M., Philippi, A., Sowa, M. E. & Lichtarge, O. (2002). Structural clusters of evolutionary trace residues are

- statistically significant and common in proteins. *J. Mol. Biol.* **316**, 139–154.
31. Thompson, J. D., Gibson, T. J., Plewniak, F., Jeanmougin, F. & Higgins, D. G. (1997). The CLUSTAL_X windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. *Nucleic Acids Res.* **25**, 4876–4882.
 32. Huelsenbeck, J. P., Ronquist, F., Nielsen, R. & Bollback, J. P. (2001). Bayesian inference of phylogeny and its impact on evolutionary biology. *Science*, **294**, 2310–2314.
 33. Ronquist, F. & Huelsenbeck, J. P. (2003). MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics*, **19**, 1572–1574.
 34. DeLano, W. L. (2002). *The PyMOL Molecular Graphics System*. DeLano Scientific, San Carlos, CA.
 35. Jones, S., Marin, A. & Thornton, J. M. (2000). Protein domain interfaces: characterization and comparison with oligomeric protein interfaces. *Protein Eng.* **13**, 77–82.
 36. Nakayama, M., Kikuno, R. & Ohara, O. (2002). Protein–protein interactions between large proteins: two-hybrid screening using a functionally classified library composed of long cDNAs. *Genome Res.* **12**, 1773–1784.
 37. Wing, M. R., Snyder, J. T., Sondek, J. & Harden, T. K. (2003). Direct activation of phospholipase C- ϵ by Rho. *J. Biol. Chem.* **278**, 41253–41258.