NON-GTPASE PROTEINS THAT SHARE COMMON MOTIFS WITH G DOMAINS: CONVERGENT OR DIVERGENT EVOLUTION OR DOMAIN RECOMBINATION?

Por

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Abstract


GTPases constitute a super class of proteins with a common fold. Five specific G motifs located in loops are signatures of this super class. Nevertheless, some proteins may share the fold of the small GTPases, although their functions are totally unrelated. To retrieve them, we specifically searched in the BLAST output listings for non GTPases with available 3D structure, starting from a canonical GTPase sequence as query. We then performed both a sequence analysis by means of HCA and a structural comparison with an established GTPase. It results that, although sequence identity is in the twilight zone, i.e. below 25%, one can evidence some conservations of the catalytic motifs. Nevertheless, mutations have occurred that produced a new function while the global fold is maintained. We discuss whether non-GTPases presumably originated from a common ancestor with an ancient G domain. The evolutive mechanisms relating non-GTPases to GTPases that we can advance are sequence divergence, convergence and DNA recombination. We conclude that the most probable evolutive pathway leading to such structural similarities is that all the studied proteins must have evolved by sequence divergence from a primordial GTP-binding domain.

Key words: HCA, GTP-binding domain, GTPase, protein evolution, protein folding, hydrophobic packing, sequence identity.

List of abbreviations: HCA, Hydrophobic Cluster Analysis; aa, amino acid(s); rmsd: root mean square deviation; ETK, E. coli protein tyrosine kinase.

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Resumen

Las GTPasas constituyen una superclase de proteínas con un plegamiento común. Cinco motivos G específicos, situados en los bucles, son característicos de esta superclase. Sin embargo, algunas proteínas adoptan el plegamiento de las GTPasas, aunque sus funciones son totalmente diferentes. Para encontrarlas, hemos analizado los resultados de búsquedas BLAST con secuencias canónicas de GTPasas, con el propósito de identificar proteínas no GTPasas con estructura 3D disponible. Posteriormente, procedimos a analizar las secuencias seleccionadas, mediante HCA y la superposición con estructuras de GTPasas de referencia. Los resultados obtenidos indican que aunque la identidad de secuencia se encuentra en la zona crepuscular (twilight zone), i.e., por debajo de 25%, se pueden evidenciar algunas conservaciones de los motivos catalíticos. Sin embargo, las mutaciones que se han producido dieron lugar a nuevas funciones, mientras que el plegamiento global se mantiene. Finalmente, discutimos si aquellas proteínas no GTPasas se originaron presumiblemente de un ancestro común con un dominio G antiguo. En tal caso, proponemos como mecanismos evolutivos que vinculan a las GTPasas con las no GTPasas, la divergencia, la convergencia y la recombinación del DNA. Concluimos que el mecanismo evolutivo más probable que dio lugar a tales similaridades estructurales es la divergencia desde un dominio primordial de unión al GTP.

Palabras clave: HCA, dominio de unión al ADN, GTPasa, evolución proteica, plegamiento de proteínas, plegamiento hidrofóbico, identidad de secuencia

1. Introducción

Small G proteins (also called small GTPases, small GTP binding proteins and Ras protein superfamily) comprise a wide variety of proteins that share the same architecture of their GTP-binding domain. Although the basic fold of this globular domain is structurally the same for all members of the family, its primary structure is extremely variable in its amino acid (aa) composition (Caldon et al., 2001). Typically, the GTP-binding domain (or G domain) is arranged in 5 α-helices (α1-α5) and six β-strands (β1-β6) (Bourne et al., 1991). Five loops named G1-G5, connecting adjacent strands and helices, contain most of the residues of the active site. Because the small GTPases are involved in diverse cellular processes (cell proliferation, protein synthesis, signal transduction), consensus sequences for the loops G1-G5 are given for each one (Paduch et al., 2001). These loops are essential for the interaction with the substrate in association with Mg²⁺ and GTP/GDP exchanging factors (Valencia et al., 1991). GTP hydrolysis “enables different GTPases to sort and amplify transmembrane signals, direct the synthesis and translocation of proteins, guide vesicular traffic through the cytoplasm, and control proliferation and differentiation of animal cells” (Bourne et al., 1991). However, various members of the GTPase superclass lack their GTPase activity (Leipe et al., 2002).

GTPases and ATPases are closely related in structure and, in fact, some ATPases constitute a subfamily of the GTPase superclass. The specificity to GTP is conferred by the NKXD motif in the G4 loop. A mutation in these aa leads to the loss of affinity for GTP, decreasing protein specificity for this substrate, but increasing for other ones. For instance, in the group of myosins and kinesins, the GTPase activity was replaced by an ATPase, because of an evolutive change in the G4 loop. However, myosin and kinesin 3D structures superpose well with that of ras proteins and are classified as a subgroup within the GTPase family (Leipe et al., 2002). The conserved Asp residue in the NKXD motif, providing specificity for GTP, is absent in most of ATPases. Thus, none of ATPases has been shown to have GTP specificity.

The aim of this paper is to provide evidence that a diverse set of proteins roughly presents the same fold as the GTP-binding domain. Nevertheless, they do not have GTPase activity and on the contrary, they exhibit activities such ribonuclease, methyltransferase, guanine deaminase, lactate dehydrogenase and others.

By means of hydrophobic cluster analysis (HCA) and comparison of structures as evidence, we discuss whether non-GTPases presumably derive from a common ancestor with an initial GTP-binding domain. Besides, the evolutive mechanisms concerning sequence divergence, convergence and DNA recombination are also considered.

2. Materials and methods

2.1. Sequence alignment and protein superposition

In order to retrieve related sequences, BLAST is used in a first step, with four proteins known from the literature
to be G domains as input data, namely: c-Ha-ras I, Gα1, RhoA and ORFX (predicted G domain-like). The output listing of BLAST provides in these four cases, a set of G domains in the top ranked sequences. We discard these first matches because they are all already known to be highly similar and we were interested in the extension of the family. Therefore, we looked at the bottom of the sorted sequences by BLAST and focused at the first sequences not annotated as G domains in the header annotations, i.e., presenting very low identity scores with the query. The retrieved sequences were aligned on short fragments (20-120 aa). In order to produce an automatic extension, alignment on the total length of the presumably unrelated sequences (typically 180 aa) was done with ClustalW online at the Pole BioInformatique Lyonnais (http://npsa-pbil.ibcp.fr/), using default parameters (Larkin et al. 2007). When comparing sequences in the twilight zone (Rost, 1999), it is known that automatic procedures do not provide accurate alignments, therefore they were manually cured by means of HCA.

2.2. Hydrophobic Cluster Analysis

HCA plots, sequence identity and HCA score calculations were performed following the indications of (Callebaut et al., 1997). Briefly, HCA designs a sequence on the surface of a cylinder with the connectivity of an alpha helix. The 2D planar surface is then duplicated in order to keep local environment for each amino acid, and hydrophobic neighbor residues (VILFMWY) in this plot are then clustered. The shapes of the clusters are keen indication of the nature of the secondary structure. Besides, it has been statistically demonstrated that centers of the hydrophobic clusters correspond to the centers of regular secondary structures (Woodcock et al., 1992). HCA identity is calculated by the number of identical aligned hydrophobic and non hydrophobic aa in both sequences to the number of aa of the longest sequence. For each sequence, the HCA score is the ratio between the number of topologically conserved residues between sequences 1 and 2, to the total number of hydrophobic residues in both segments (Gaboriaud et al., 1987). A HCA score ≥ 60% is an indicator of high sequence identity.

2.3. Structural comparisons

As evolution has more conserved structures than sequences (Grishin, 2001; Wang et al., 2008), in a final step, structure superpositions were performed to validate our previous hypotheses. Tertiary structures were all obtained from the PDB (Berman et al., 2008) and their superpositions were carried out with the on line programs MATRAS (Kawabata, 2003) and CE (Shindyalov & Bourne, 1998). Superpositions were performed without forcing the structural alignment to match the sequence alignment derived from HCA. Matras and CE produced similar results. Thus, Matras has been chosen, because we noticed in previous papers that its results are the closest from a consensus of three methods (Papandreou et al., 2004).

Except step 2, which is a specialty of the Paris group, the work flow used here is rather classical: an identification of a hot spot in a short stretch of sequences, followed by a structural validation. This has been done for instance on the short chain oxidoreductase enzymes (Duax et al., 2007), SH3 fold (Theobald & Wuttke, 2005) or on more general sequences (Liu et al., 2008). The scheme is the following: first sequences are compared in order to derive evidence for a common ancestry. In case of absence of similarity, structure similarity is search for.

3. Results

3.1 Protein sequence database search

When performing BLAST searches with a GTP-binding domain, it is usual to find non-GTPase proteins that share 50-100 aa long fragments with G domains, with a level of identities located in the “twilight zone” of sequence alignment (Rost, 1999). Subsequent superpositions of their corresponding 3D structures yield rmsd lower than 6 Å (Reva et al., 1998), that can be considered as a reasonable limit to admit that they share a common fold, and therefore are related through evolution. Thus, we were interested to elucidate the evolutive relationships among these proteins and the molecular mechanisms leading to the gain of a similar architecture.

A database screening was carried out with G domain sequences, in order to retrieve non-GTPase proteins with both low 1D identity and partial 3D similarity, and analyze them by HCA analysis and protein superposition. The purpose of such alignments was to elucidate the most conserved catalytic and structural amino acids with the G domain. HCA is a fine method of 2D structural analysis that allows alignments between very distantly related proteins, with as low as 10% sequence identity. HCA analysis does not pay any attention to the strict conservation of the residues inside the clusters but rather to the conserved shapes of the clusters, keeping in mind the underlying idea that shape is a testimony of the secondary structure. As shown as follows, HCA was an outstanding method to reveal structural similarities of proteins like GTPases and non-GTPases, which at first glance do not appear to be related.
BLAST screening of PDB (Berman et al., 2008) and Blocks (Henikoff & Henikoff, 1994) databases was done with the following G domain primary structures as queries: (c-Ha-ras1 (HRas precursor, P01112), Guanine nucleotide-binding protein G\text{\alpha} \text{subunit} (G\text{\alpha}, P04898) and ORFX (AF261774)). ORFX is a protein of unknown function from \textit{Lyceopersicon esculentum} that is involved in determination of fruit size and seems to have a predicted fold similar to the one of ras (Frary et al., 2000).

Most of the retrieved sequences were GTP-binding domains, with high identity with the query domains. However, a few sequences of non-GTPase proteins exhibited local identities in the range of 10 to 23% with G domains. Then, we selected the first non-GTPase sequences in the list for primary, secondary and tertiary sequence alignments by means of ClustalW, HCA and 3D structure superposition, respectively. We did not find sequences that meet our criteria in other than PDB and Blocks databases. In Table 1, we list the retrieved sequences. HCA identities were always slightly higher than ClustalW identities in the range 18 to 30%. This is actually frequent, because HCA produces a more sensitive alignment than a standard program. HCA scores were in the range 55-60%, a good indicator of a significant homology (Gaboriaud et al., 1987). Low rmsd values are also listed in Table 1, following structure superposition.

To illustrate the use of HCA alignments in combination with protein superpositions and the qualitative and quantitative information we can derive from such alignments, we show an example in Fig. 1 and Table 1. Two pairs of distantly related proteins with low 1D identity, the human c-Ha-ras1 (a GTPase) with RhoA (a GTPase) (Gómez del Pulgar et al., 2005) and c-Ha-ras1 (a GTPase) with the CMP kinase (an ATPase) (1Q3T), were aligned by HCA. Previously, a 35% aa identity has been evidenced between rho genes from \textit{Aplysia} and c-Ha-ras1 (Madaule and Axel, 1985). As seen in Fig. 1a GTPases share most of the hydrophobic clusters (1D=29%) and 3D structures superpose along the entire domain (rmsd 1.62 Å). However, the GTPase and the ATPase (Fig. 1b), which aroused from a unique primordial fold (Leipe et al., 2003), share an identity of 12% (increased to 23% by HCA) and partial superposing subdomains: the N-terminal 32 aa (i.e., G1 motif and some aa of G2) produces a rmsd of 5.4 Å and the C-terminal 45 aa (G4 and G5 motifs), a rmsd of 5.1 Å. Interestingly, the G3 loop (the so-called Walker B box (Walker et al., 1982) or kinase 2 motif) conserves both hydrophobic clusters and identities in the GTPase and the ATPase proteins, although they are located at the end of a β-sheet in c-Ha-ras1 and a \(\alpha\)-helix in CMP kinase. The G3 loop provides residues for Mg\textsuperscript{2+} and γ-phosphate binding and is found in other nucleotide binding motifs, not homologous to small G proteins, present in sugar kinases, ABC transporters and ATP synthases (Paduch et al., 2001). Thus, despite the degree of divergence between GTPases and related ATPases, structural relationships can be evidenced by a refined alignment method like HCA and supported by structure superposition. Our data confirm the tight structural relation of the GTPase and ATPase, notwithstanding evolutive divergences.

### Table 1. Sequence identities among pairs of proteins, after domain alignments performed with ClustalW or HCA. The approximate length (in amino acids) along which the alignment is performed is indicated in the aa column. HCA scores were calculated as stated by Gaboriaud et al., (1987). Rmsd over superposed length values (number of aa) are shown. In most of the cases the superposition is performed on a whole domain, but in two cases, corresponding to 1b and 3a, a hinge fragment is evidenced from the HCA analysis. Consequently, the superpositions have been calculated on the parts surrounding these hinge regions. Uniprot accession numbers are the following: P01112 for c-Ha-Ras; P61186 for RhoA; Q97PK6 for CMP kinase; P04898 for G\text{\alpha}; Q57599 for Rnase HII; Q9GT92 for LDH; P55135 for Rum A; O34598 for Guanine deaminase; af261774 for OrfX.

<table>
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<tr>
<th>Figure</th>
<th>Aligned proteins</th>
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<th>HCA</th>
<th>HCA Score</th>
<th>aa</th>
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<td>30</td>
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<td>23</td>
<td>59</td>
<td>170</td>
<td>5.4/32; 5.1/45</td>
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<tr>
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<td>G\text{\alpha} - RNase HII</td>
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<td>20</td>
<td>56</td>
<td>170</td>
<td>4.03/79</td>
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<tr>
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<td>RhoA - LDH</td>
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<td>22</td>
<td>62</td>
<td>170</td>
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<tr>
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<td>170</td>
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Figure 1. HCA plots of pairs of aligned sequences. a) Two low identity GTPases, c-Ha-ras\textsubscript{1} (PDB code 4Q21) and RhoA (PDB code 1X86\textsubscript{b}); b) c-Ha-ras\textsubscript{1} aligned with the CMP kinase (PDB code 1Q3T, ATPase). 

Symbols: Conserved hydrophobic clusters are grey shaded. Non hydrophobic identities are indicated by white letters inside black circles. White letters on a grey square indicate catalytic aa in G motifs (G1-G5 on top) and residue conservation in non-GTPases. Black or white aa in a larger font size (some inside black circles) are residues that interact with a specific ligand as cofactors or substrates. Under each plot, the secondary structures are schemed, based on PDB files. The boxed regions indicate the fragments of the proteins for which the structures can be superimposed. The onset helps interpreting the HCA plots. Because of the duplication (see methods), sequence is read vertically, one line over two, and the secondary structure is read horizontally, a cluster corresponding statistically to a regular secondary structure. Vertical lines connect the occurrences of analogous clusters.
methyluridine (specifically to uridine 1939 of 23S rRNA to yield 5- of a methyl group from S-adenosylmethionine (SAM), (Q5JHF7). In coenzyme NADH. With GTP have evolved to be able to bind the dinucleotide the two proteins. Hence, the aa pertinent for interaction to conclude to the existence of a partial common fold for positions occupied by catalytic residues once more allow ItN (120-122) (conserved residues in capital letters) in G4 (positions 14-15) are in the G1 motif of RhoA, D35 in G2, are located around G1-G5 motifs. For example, residues QI (NADH). The aa involved in the interaction with the ligand complexed with substrate (pyruvic acid) and cofactor and G5 motifs. The LDH structure (PDB code 2FM3) is 3.02 Å over 78 aa in the C-terminal region, including the G4 superposition of the two domains, we obtained a rmsd of 4.03 Å over 79 aa at the C-terminal end. A striking observation in the Archaeal RNase HII of the hypothermophile Methanococcus jannaschii (Lai et al., 2000), homologous to the human major RNase H. ClustalW and HCA identities were 14 and 20%, respectively. In Fig. 2a we can observe that the most relevant conserved hydrophobic clusters are mainly associated to ǂ-helices, which are rather long for RNase HII. Structure superposition gave a rmsd of 4.03 Å over 79 aa at the C-terminal end. A striking observation in Fig. 2a is that the most conserved secondary structures and catalytic residues are in boundary of G4 and G5 loops, and G4 is the motif that determines the interaction with GTP or ATP (see introduction). Because RNase HII interacts with a polymer of nucleotides (E. coli RNase HII cleaves the RNA strand of a RNA-DNA hybrid, endonucleolytically at the P-O3’ bond), one can speculate that the motifs responsible for nucleotide interaction and hydrolysis have been rearranged to perform RNA degradation.

When using RhoA as a query sequence, we obtained the lactate dehydrogenase (LDH) from Cryptosporidium parvum. Sequence identities were 18 and 22% for ClustalW and HCA alignments, respectively (Fig. 2b). After superposition of the two domains, we obtained a rmsd of 3.02 Å over 78 aa in the C-terminal region, including the G4 and G5 motifs. The LDH structure (PDB code 2FM3) is complexed with substrate (pyruvic acid) and cofactor (NADH). The aa involved in the interaction with the ligand are located around G1-G5 motifs. For example, residues QI (positions 14-15) are in the G1 motif of RhoA, D35 in G2, ItN (120-122) (conserved residues in capital letters) in G4 and MagV (145-148) in G5. Thus the similar distribution of hydrophobic clusters, the low rmsd and the coincident positions occupied by catalytic residues once more allow to conclude to the existence of a partial common fold for the two proteins. Hence, the aa pertinent for interaction with GTP have evolved to be able to bind the dinucleotide coenzyme NADH.

With ORFX (see below) as BLAST query, we retrieved in Blocks database the Block number IPB010280C (motifs G4-G5) of Thermococcus kodakarenensis RumA protein (Q5JHF7). In E. coli, this enzyme that catalyzes the transfer of a methyl group from S-adenosylmethionine (SAM), specifically to uridine 1939 of 23S rRNA to yield 5'-methyluridine (Lee et al., 2004). We show in Fig. 2c a HCA alignment between E. coli RumA and c-Ha-ras1, not ORFX, because of the absence of PDB structures; however ORFX is predicted to superpose well with c-Ha-ras1 since their Z scores for global and local alignments are high (3.2 and 4, respectively) (Frary et al., 2000). Identities are 10% and 18% with ClustalW and HCA, respectively. The major cluster and secondary structure conservations are located, like in RNase HII and LDH, at the C-terminal end (Fig. 2c). We were able to superpose these regions, obtaining a rmsd of 3.46 Å over 64 aa. Lee et al. locate the putative site of interaction of RumA with SAM to residues 268-282 (AGV…EWL, Fig. 2c) (Lee et al., 2004). These positions perfectly match with G1 and G2 motifs in c-Ha-ras1.

With c-Ha-ras1 as a query sequence, we retrieved the Bacillus subtilis Guanine Deaminase, an enzyme that catalyzes the hydrolytic deamination of guanine into xanthine (Liau et al., 2004) (Fig. 3a). Sequence identities are 10 and 18%. The high conservation of secondary structures and their direct relation with hydrophobic clusters in number and shape are obvious in Fig. 3a. A protein superposition was possible in two fragments (Fig. 3a, Table 1) yielding a rmsd of 3.87 Å over 28 aa (N-terminal) and 4.27Å over 55 aa (middle of the protein). 1-2 aa in each G1-G4 motifs are shared between the two sequences.

The last sequence we analyzed is ORFX (AF261774), one of the most intriguing proteins with a probable G-domain fold. It is a protein of unknown function, but involved in determination of fruit size in tomato (Frary et al., 2000) and classified in Pfam as a member of the PLAC8 family (Placenta-specific gene 8 protein). ORFX was predicted in the literature as sharing a similar fold (LOOPP program (Tobi & Elber, 2000) than the human oncogene ras protein (PDB code 6Q21). We show in Fig. 3b a HCA alignment of c-Ha-ras1 and ORFX. A striking observation is the high correspondence of hydrophobic clusters and aa conservation of the G motifs, excepting G3, with a HCA identity of 17% (Table I). Unfortunately there is no available 3D structure for protein superposition. Because of its assumed regulatory activity, we tried to associate ORFX with transcription factors. HCA analysis shows that ORFX is enriched in cysteines, a common characteristic of zinc finger proteins; thus, we aligned ORFX with members of the zinc finger family and we show an example with Atzfp6 (Swiss-Prot entry Q39265) in Fig. 3c. Interestingly, the numerous cysteines of ORFX are in the same domain organization than zinc finger proteins (ZF1 and ZF2 in Fig. 3c). A prediction of zinc-binding cysteines with Predzinc (Shu et al., 2008) revealed four candidate residues with probability scores exceptionally high to constitute a putative zinc-binding domain (ZF2, Fig. 3c): Asp111 (0.576), Cys118 (0.742), Cys121 (0.742) and Cys124 (0.656). Besides,
Figure 2. HCA plots of pairs of aligned sequences. a) $G_{ia}$ (PDB code 2EBC, GTPase) aligned with RNase HII (PDB code 1EKE, non GTPase); b) RhoA (PDB code 1X86_B, GTPase), aligned with LDH (PDB code 2FM3, non GTPase); c) c-Ha-ras1 (PDB code 4Q21, GTPase), RumA (PDB code 1UWV, non GTPase). The HSSP producing hits with BLAST are marked by a black line under the alignments.
Figure 3. HCA plots of pairs of aligned sequences. a) c-Ha-ras1 (PDB code 4Q21, GTPase) aligned with guanine deaminase (PDB code 1WKQ, non GTPase); b) c-Ha-ras1 aligned with ORFX c) ORFX (accession number af261774) aligned with Atzfp6 (Q39265).
we can observe in Fig. 3c the remarkable residue conservation of hydrophobic (clusters) and non-hydrophobic residues for motifs 1 (23% identity), 2 (20% identity) and 3. These results enable us to propose that ORFX may originate from a primordial G domain and belong to an unknown class of zinc finger proteins with transcription regulatory functions.

In order to supply a graphic view of our hypothesis of inclusion of new members in the G domain family, we show in Fig. 4 structural superpositions of RhoA with LDH on top, and c-Has-ras1 superposed on RumA at the bottom. Global rmsd in these two cases is 3.02 Å and 3.46 Å, respectively, therefore validating our assumption of two new members in the family.

![Figure 4. Superposed stereo structures of a protein known to belong to the G domain (RhoA (P^71-G^155) on top and c-Has-ras1 (R^81-S^144) at the bottom) with a proposed new member of the family (LDH (D^84-C^161) on top and RumA (Q^350-L^414) at the bottom.](image)

4. Discussion

The phylogenetic tree of GTPase and ATPase families is now well established and in numerous cases they originate from a common nucleotide hydrolytic domain (Leipe et al., 2002; Leipe et al., 2003). However, some proteins share very low identity with the other members of the superclass and are difficult to classify in this family by means of sequence alignments. Structure superposition is of great help to produce a reliable alignment. A decisive criterion to incorporate a putative GTPase in the family is the conservation of the G1 to G5 motifs. Even though G domains may be highly divergent, they are still recognizable by the appropriate methods (Fig. 1).
these proteins are nucleotides or polymers of nucleotides; thus, the structural affinity for nucleotides is maintained but with novel activities. v) Previously, we have demonstrated that a GTP-binding domain was recycled to have a receptor activity in the A domain of the chloroplast GTPase receptor Toc159 (Hernández Torres et al., 2007). This is one example of adaptation of the G domain to a new function, independent of nucleotide binding. A second case of a GTPase domain adaptation is ORFX. As seen in Fig. 3b, the alignment between ras and ORFX proteins is well consistent in hydrophobic clusters, as well as in non-hydrophobic aa (17% identity). In this work we propose that ORFX has a function of a zinc-finger transcription factor and we present strong evidence in Fig. 3c. It is no clear how a nucleotide hydrolyzing domain becomes a transcription factor; however its primary nucleotide binding activity and the flexibility of loop regions must have been the key for this domain transformation (Blouin et al., 2004).

A second way by which non-GTPase proteins may have appeared is convergent molecular evolution (CME). There are few reports demonstrating this evolutive event, somehow controversial. CME is defined as “the independent evolution of similar nucleotide or amino acid sequences in two unrelated molecules” (Zakon, 2002). Structural convergence is one of the four classes of CME and could be the most suitable for non-GTPase proteins. As stated by Zakon (2002), “in some cases, molecules with very different amino acid sequences can assume similar structural motifs, and these may carry out similar functions”. If it was the case for non-GTPase proteins, the convergence would be only in the 3D structure but not in the functional level. The best models of reported CME refer to different structures with the same function (see for example the hexokinase, ribokinase, and galactokinase families of sugar kinases (Bork et al., 1993). There are two points that refute a case of CME for non GTPases: i) convergence would occur only for structure but not for function and ii) all the aligned proteins display conserved functional residues with GTPase G1-G5 motifs (Figs. 2 and 3). It would be difficult to accept all those analogous residues as a product of chance, particularly if each protein exhibits a different metabolic function.

The third probable mechanism at the origin of non-GTPases could be the partial recombination between ancient genes and a primordial GTP-binding domain, in order to produce new genes which gained only some of G1-G5 motifs. However, this alternative would be less probable than convergence, because the high conservation of hydrophobic and identity clusters, as well as functional residues.

Taking together all results, we conclude that the most feasible mechanism by which non-GTPases hold reminiscences of a common ancestral GTP-binding domain is divergence. Then it was simpler for nature to assign new functions to an already existing nucleotide-interacting domain, with a particularly flexible architecture in their catalytic loops, than building new ones from scratch.

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