Research article

Novel conserved domains in proteins with predicted roles in eukaryotic cell-cycle regulation, decapping and RNA stability

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Abstract

Background: The emergence of eukaryotes was characterized by the expansion and diversification of several ancient RNA-binding domains and the apparent de novo innovation of new RNA-binding domains. The identification of these RNA-binding domains may throw light on the emergence of eukaryote-specific systems of RNA metabolism.

Results: Using sensitive sequence profile searches, homology-based fold recognition and sequence-structure superpositions, we identified novel, divergent versions of the Sm domain in the Scd6p family of proteins. This family of Sm-related domains shares certain features of conventional Sm domains, which are required for binding RNA, in addition to possessing some unique conserved features. We also show that these proteins contain a second previously uncharacterized C-terminal domain, termed the FDF domain (after a conserved sequence motif in this domain). The FDF domain is also found in the fungal Dcp3p-like and the animal FLJ22128-like proteins, where it fused to a C-terminal domain of the YjeF-N domain family. In addition to the FDF domains, the FLJ22128-like proteins contain yet another divergent version of the Sm domain at their extreme N-terminus. We show that the YjeF-N domains represent a novel version of the Rossmann fold that has acquired a set of catalytic residues and structural features that distinguish them from the conventional dehydrogenases.

Conclusions: Several lines of contextual information suggest that the Scd6p family and the Dcp3p-like proteins are conserved components of the eukaryotic RNA metabolism system. We propose that the novel domains reported here, namely the divergent versions of the Sm domain and the FDF domain may mediate specific RNA-protein and protein-protein interactions in cytoplasmic ribonucleoprotein complexes. More specifically, the protein complexes containing Sm-like domains of the Scd6p family are predicted to regulate the stability of mRNA encoding proteins involved in cell cycle progression and vesicular assembly. The Dcp3p and FLJ22128 proteins may localize to the cytoplasmic processing bodies and possibly catalyze a specific processing step in the decapping pathway. The explosive diversification of Sm domains appears to have played a role in the emergence of several uniquely eukaryotic ribonucleoprotein complexes, including those involved in decapping and mRNA stability.
Background

Systematic comparative analyses of genome sequences have suggested that the majority of domains found in proteins involved in RNA metabolism are drawn from a relatively small set of conserved domains (approximately 100–135) [1-3]. The proteins containing these conserved domains correspond to around 4 to 11 percent of the protein-coding genes in cellular life forms and perform a wide range of functions that include translation and its regulation, processing and modification of cellular RNAs, and post-transcriptional gene regulation [1-3]. This set of conserved domains can be broadly divided into those that mediate interactions with RNAs or other proteins in ribonucleoprotein complexes, and catalytic domains that may catalyze a wide range of reactions related to RNA or associated proteins. Most of the common RNA-binding domains (RBDs) are relatively small (less than 150 residues) and tend to be evolutionarily mobile, occurring as solos, or in combination with other RBDs or enzymatic domains [1]. Several RBDs as well as the catalytic domains of RNA metabolism enzymes are amongst the most highly conserved and universally distributed protein domains in cellular organisms. These highly conserved domains are typically present in ribosomal components, translation factors, enzymes that modify RNA and tRNA, polyadenylation, and transcription elongation factors [1,4,5]. However, the analysis of phyletic patterns of conserved domains has also suggested that a significant innovation of novel RBDs occurred at the base of eukaryotes [1]. These eukaryotic innovations include the PAZ, G-Patch, PWI and SWAP domains and several Zn-chelating domains, such as the Zn-knuckle, the CCCH and LRP fingers [1,6-8]. The emergence of these domains, as well as the expansion and diversification of superfamilies of previously existing domains appears to have accompanied development of several novel aspects of RNA metabolism in the eukaryotes. These unique eukaryotic aspects include pathways involved in pre-mRNA splicing, capping, post-transcriptional gene silencing and nucleo-cytoplasmic RNA transport. The eukaryotes also possess more complex versions of RNA degradation and processing systems, such as the exosome and the multi-subunit RNaseP/RNase MRP [1,9,10]. Hence, the identification of novel eukaryote-specific domains, as well as the analysis of the diversification of ancient domain superfamilies in eukaryotes may help in providing a better understanding of the origins and the biochemical properties of the unique aspects their RNA metabolism.

The computational identification of conserved RNA-binding domains (RBDs) has considerably contributed to the analysis of RNA-protein interactions in various pathways of RNA metabolism [1,6,11-13]. The enzymatic domains associated with RNA metabolism typically belong to superfamilies, which may also include members that act on substrates outside the context of RNA metabolism (eg. Rossmann fold methyltransferases acting on non-ribonucleoprotein substrates) [1]. Hence, the combinations of RBDs and enzymatic domains in the same polypeptide provide a strong contextual handle for predicting novel catalytic activities associated with RNA metabolism. Comprehensive analysis of the commonly occurring domains involved in RNA metabolism has previously helped in identifying several such domain architectures that led to the prediction of novel RNA and RNP modifying/processing enzymes [1,6,14,15]. The recent increase in the available genomic sequences from eukaryotes provides further opportunities to extract contextual information in the form of previously unnoticed domain architectures. Furthermore, the new data also allows the detection of less common, nevertheless functionally important eukaryote-specific domains, which may have eluded earlier screens for such domains. Additionally, other forms of contextual information emerging from newer studies involving large-scale mutational analysis of eukaryotic genes, high-throughput analysis of gene expression, sub-cellular protein localization and protein-protein interactions could also provide clues regarding the functions of uncharacterized proteins.

In particular, we are interested in using computational methods to identify novel eukaryote-specific proteins that may be involved in RNA metabolism and predicting their potential biochemical functions. In the current work we use a combination of sequence analysis, homology-based fold prediction and contextual information to describe two novel conserved RNA-protein or protein-protein interaction modules and one catalytic module that are found in proteins predicted to participate in regulation of the cell cycle and decapping. We discuss these findings in the context of the origin of the decapping apparatus in eukaryotes and present hypotheses for the possible functions of poorly characterized but highly conserved groups of eukaryotic proteins.

Results and discussion

Identification of the novel FDF domain and conserved eukaryotic proteins with domains related to the RNA-binding domain SM domain

Several RNA binding proteins in eukaryotes are characterized by the presence of highly charged or polar low-complexity segments, typically containing repeats of simple motifs such as SR, RG and GGY [16-18]. Experimental evidence has suggested that these segments interact with RNA with low target specificity or aid in their localization to specific RNA processing substructures [16-19]. These segments are usually combined with globular domains that may mediate more specific interactions with RNA. Hence, detection of proteins containing these segments provides a means of identifying potential RNA-binding
proteins that may either lack previously characterized RBDs or contain very divergent versions of them. Accordingly, we generated a sieve for such proteins using pattern searches that identified proteins with multiple occurrences of the low-entropy repeat motifs that are typical of RNA-binding proteins. Those proteins in this set, which were identified as potential RNA-binding proteins or RNA-processing enzymes in our previous surveys [1,20,21] conducted using sensitive profiles for RBDs and associated enzymes, were removed in the first step. Of the proteins that remained, we selected those proteins that contained potential globular domains when screened using the SEG program [22]. These proteins were then further searched using the Pfam domain collection [23] to identify any previously reported modules that may have escaped our searches.

Via this procedure we identified one group of experimentally uncharacterized proteins typified by Saccharomyces cerevisiae Scd6p and Schizosaccharomyces pombe Sum2p as potential RNA-binding proteins. These proteins formed a distinctive family (hereinafter Scd6p family), which included the mRNA binding protein Rap55 from the newt Pleurodeles waltl and orthologous representatives from fungi, animals, plants and apicomplexans (Cryptosporidium and Plasmodium). This observation suggests that the family is likely to have emerged prior to the diversification of the crown group eukaryotes and possibly performs a well-conserved function. Analysis with the SEG program [22] suggested that these proteins contain distinct N- and C-terminal globular domains flanked by low complexity regions enriched in charged residues, including the RS and RG motifs. In order to understand better the affinities of these globular domains we initiated PSI-BLAST searches (profile inclusion threshold = .01; iterated to convergence) of the Non-Redundant database (NR) with them using representatives from several different organisms. Interestingly, in searches with the N-terminal module, Sm RNA-binding domains were recovered, either with significant hits ($e = 10^{-4}$–$10^{-6}$) or as the best hits with borderline E-values. As these domains had not been reported by others or us in systematic surveys for Sm proteins [1,24], we investigated them in greater detail using new position-specific score matrices, which were made by including all the previously identified representatives of Sm domains in the nr database. A search of the NR database with this profile recovered members of the Scd6p in iteration 7 with significant e-values ($e = 10^{-4}$–$10^{-6}$ at the point of first recovery). Secondary structure prediction using a multiple alignment of the of the N-terminal globular domain of the Scd6p family showed that it possessed an all β-fold with a perfect correspondence to the secondary structure elements observed in the Sm-type SH3 β-barrel fold [25,26] (also see SCOP Database [27]). Barring the Sm domains, neither other members of the SH3-like folds nor any other distinct β-barrel-folds, such as the OB fold, were recovered in these searches. Likewise, the Scd6p-like proteins were not detected in searches with profiles for various OB fold domains and other β-strand rich RNA-binding domains. These observations strongly suggested that the Scd6p family contained a previously unreported, divergent form of the Sm domain.

A multiple alignment of the classical Sm domain was generated using a structural superposition of all crystallized Sm domains proteins from the PDB database, including the divergent bacterial version Hfq, as a template (Fig. 1). A comparison of the multiple alignment of the Scd6p family with this alignment of Sm domains shows that it contains the hallmark features of the latter class, such as the presence of a hxG signature (where h is a hydrophobic residue) in the N-terminal half and a +Gpph signature (where `p' is a polar residue and `+' a positively charged residue), which is seen in the C-terminal half of the archaeo-eukaryotic versions (Fig. 1). Additionally the Scd6p family contains certain unique features that set it apart from other Sm domains: 1) It contains a conserved C-terminal extension that is likely to form an additional terminal strand that is usually lacking in many of the classical Sm domains (Fig. 1). 2) It contains a characteristic motif, usually of the form GTEx+ (where + is a positively charged residue; x is any residue) in the variable region separating the conserved N- and C-terminal halves of the Sm domain (Fig. 1). Most Sm domains contain a helix of variable length at their N-terminus [28]. The Scd6p family shows relatively poor sequence conservation and weak helix prediction in the corresponding N-terminal regions. However, the presence of the conservation in the Scd6p family of the capping residue (either glycine or a small residue), which is present in the C-terminus of this helix, suggests that it might contain an abbreviated version of this helix (Fig. 1).

The Sm proteins from archaea and eukaryotes and the bacterial Hfq proteins do not bind RNAs stably as monomers, but only as heptameric or hexameric toroids [25,28]. Furthermore, even the highly divergent versions of the Sm superfamily, such as the MscS protein of the bacterial mechano-sensory channels [29], form heptameric toroids similar to the RNA binding Sm domains, suggesting that this quaternary structure may be pervasive throughout this superfamily. Accordingly, we speculate that the Scd6p proteins are also likely to be incorporated into such structures. When the conservation pattern of the Scd6p proteins is compared to the RNA contacts of the Sm domains in the crystal structure of the Archaeoglobus fulgidus Sm1 (AF0875) heptameric ring, several similarities and a few notable differences are seen [25] (Fig. 1). In the highly conserved C-terminal +Gpph motif, the side chain of the positively charged residue (R63 in Af Sm1/AF0875)
Figure 1
Multiple alignment of the Scd6 family with representatives of other Sm domains. Multiple sequence alignment of the Sm domain of the Scd6 family was constructed using T-Coffee after parsing high-scoring pairs from PSI-BLAST search results. The secondary structure from the crystal structures is shown above the alignment with E representing a strand. The 90% consensus shown below the alignment was derived using the following amino acid classes: hydrophobic (h: ALICVFMYFW, yellow shading) and its aliphatic subset (l: ALIV, yellow shading); small (s: ACDBNPSTV, green); and polar (p: CDEHKNQRST, blue). The limits of the domains are indicated by the residue positions, on each end of the sequence. A '*' denotes the end of the protein sequence. The numbers within the alignment are non-conserved inserts that have not been shown. The conserved GTEx+ motif of the scd6 family is shaded red. The residues involved in RNA binding are denoted by 'HY's on the top of the alignment. The conserved C-terminal extension of the Scd6 family is shown in a box. The sequences are denoted by their gene name followed by the species abbreviation and GenBank Identifier (gi). The species abbreviations are: Af – Archaeoglobus fulgidus; Ec – Escherichia coli; Sa – Staphylococcus aureus; Afum – Aspergillus fumigatus; At – Arabidopsis thaliana; Cb – Caenorhabditis briggsae; Ce – Caenorhabditis elegans; Dm – Drosophila melanogaster; Hs – Homo sapiens; Nc – Neurospora crassa; Pf – Plasmodium falciparum; Pw – Pleuro德子 waltz; Sc – Saccharomyces cerevisiae; and Sp – Schizosaccharomyces pombe.

While the hydroxyl group of this residue might form a hydrogen bond with the base, it is unclear if it could confer the uracil-specificity that is provided by the asparagine in the canonical Sm domains. The Scd6 family has a polar residue instead of the aromatic residue that stacks against the base in most other canonical Sm domains (H37 in Af Sm1/AFO875; Fig. 1). This polar residue is likely to form hydrogen bonds with base rather than the stacking interactions which are observed in most other Sm domains [25]. These differences, along with the Scd6 family-specific GTEx+ motif that occurs between the N-
and C-terminal conserved regions of the domain, are likely to confer certain unique nucleic-acid-binding properties on the Scd6 family [30].

Most members of the Scd6 family contain a single Sm-related N-terminal domain fused to another conserved C-terminal domain, except At4g19330 from Arabidopsis, which is comprised of just two tandem repeats of the Sm domain (Fig. 2). In order to investigate the distinct C-terminal domain of the Scd6 family we initiated PSI-BLAST searches with this domain. In addition to members of the Scd6 family, these searches also recovered other proteins with significant e-values such as the Dcp3p (Yel015wp) protein from S. cerevisiae and its fungal relatives and uncharacterized proteins such as FLJ21128 (gi: 19923613) from Homo sapiens and its relatives from various animal clades. For example, searches with C-terminal domain of the human Scd6p ortholog (gi: 13559033) recovered Yel015wp/Dcp3 in iteration 5 with e = 0.003 and FLJ21128, e = 4 x 10^-4. Reciprocal searches with this region from the above-mentioned proteins, such as Dcp3p and FLJ21128 recovered bona fide members of the Scd6p family with significant e-values (e.g. the region from FLJ21128 recovered the Rap55 in iteration 3; e = 2 x 10^-4). Unlike the Scd6p family, this conserved region occurred in the N-terminal region of the Dcp3p and FLJ21128 proteins. These latter proteins additionally contained a C-terminal globular domain, which belongs of a specialized family of Rossmann fold domains. This family of Rossmann fold domains also includes the N-terminal domain of the E. coli YjeF protein and, hereinafter we refer to this domain as the YjeF-N type Rossmann fold domains (see below for further discussion).

The above observations indicated that the conserved region shared by the Scd6p family, Yel015wp/Dcp3p and FLJ21128 is likely to define a novel domain. We named it the FDF domain after the characteristic signature that is present at N-termini of these domains (Fig. 3). The multiple alignment of the FDF domain shows that it is enriched in polar and charged residues with few hydrophobic residues embedded in their midst. It is predicted to adopt an entirely α-helical structure with multiple exposed hydrophilic loops. These features suggest that the FDF domain is likely to interact with RNA or highly charged peptides that are commonly found in the ribonucleoprotein complexes. Though the animal FLJ21128-like proteins and the fungal Yel015wp/Dcp3p differ in their architectures and are considerably divergent in terms of sequence, the presence of a shared architectural core (FDF domain fused to a YjeF-N-like Rossmann fold domain), which is not found in any other eukaryotic proteins suggests that they might belong to the same orthologous lineage shared by animals and fungi (Fig. 2 and 3).

N-terminal to the FDF domain, the FLJ21128-like proteins from animals, but not the fungal Dcp3p-like proteins, contain an additional small conserved globular domain. Based on its predicted secondary structure it is likely to adopt an all β-fold. Further analysis of this globular domain using profiles for conserved domains showed that it gave a significant hit (e-value=.005–0.001) with the Sm domain profile. This observation, taken together with its conservation pattern suggests that the extreme N-terminal domain in the FLJ21128-like proteins is yet another uncharacterized, divergent version of the Sm fold (Fig. 1 and 2).

Potential functions for the FDF and Scd6p-like Sm domain proteins in cell-cycle regulation and decapping

Genetic studies on S. cerevisiae Sdc6p and S. pombe Sum2p have been fairly opaque with regards to their functions. The Scd6p has been recovered as a suppressor of clathrin deficiency [31]. However, there is no evidence that it directly functions in the assembly of clathrin-coated vesicle. High-throughput localization studies have indicated
Figure 3
A multiple sequence alignment of the FDF domain. Multiple sequence alignment of the FDF domain was constructed as described in Figure 1. In the secondary structure H represents a helix. The species abbreviations are as given in Figure 1 and additionally: Ani – Aspergillus nidulans; Gze – Gibberella zeae;Mgr – Magnaporthe grisea.

that it is localized to the cytoplasm and not the nucleus in S. cerevisiae [32]. Sum2p was recovered as a weak suppresor of the over-production of the G2/M checkpoint regulator, Cdc25p [31]. The Cdc25p phosphatase is an activator of the cyclin dependent kinase Cdk2p and when over-produced it results in a bypass of the G2/M checkpoint, which ensures that DNA replication is completed before the M phase is initiated. Specifically, expression of the N-terminal Sm-like domain of Sum2p, but not the full length Sum2p, was found to restore the G2/M checkpoint bypass in Cdc25p-overproducing cells, as well as in cells with mutations in Cdk2p and Wee1p, which show identi-

Figure 1. In the secondary structure H represents a helix. The species abbreviations are as given in Figure 1 and additionally: Ani – Aspergillus nidulans; Gze – Gibberella zeae; Mgr – Magnaporthe grisea.
production of Cdc25p in S. pombe is the Sum3 gene, which encodes a RNA helicase [31]. Hence, it is possible that Sum2p and Sum3p act together to regulate the stability and translation of a similar set of mRNAs encoding check point proteins.

The available evidence also implicates the Dcp3p and FLJ21128 proteins with FDF and YjeF-N-type Rossmann fold domains in the decapping process. High-throughput analyses of protein-protein interactions in yeast using affinity precipitation and two-hybrid systems have consistently recovered the decapping enzymes Dcp1p and Dcp2, Dhhlp, the superfamily II helicase involved in decapping process, and the ribosomal protein S28 as potential interaction partners of Dcp3p [42-44]. The subcellular localization pattern of Dcp3p based on GFP tag analysis indicates that it is entirely cytoplasmic like Scd6p, Dhhlp. Specifically, it translocates to punctate foci [32], just like the decapping enzymes Dcp1p and Dcp2p and the Lsm1-7p complex [40,45]. These observations suggest that the Dcp3p and FLJ21128 proteins are likely to be associated with other proteins of the mRNA decapping complex in the specialized cytoplasmic processing bodies [45]. The presence of the N-terminal Sm domain in the FLJ21128 (and it orthologs from other animals) suggests that it might directly interact with other Sm proteins to be incorporated in specialized Sm heptamers.

Further clues regarding the functions of the Dcp3p and FLJ21128 are furnished by an analysis of the C-terminal YjeF-N-type Rossmann fold domain. Both iterative sequence searches with the PSI-BLAST program and structural similarity searches of PDB show that the dehydrogenase-type Rossmann domains are their closest relatives. For example a PSI-BLAST search with the YjeF-N domain of Dcp3p recovers dehydrogenases with significant e-values (e = 10^-5; iteration 6), while Ynl200cp (PDB:1jzt), a member of this family, recovers oxidoreductases like D-glyceraldehyde dehydrogenase with significant Z-scores (Z = 8.9) in structural similarity searches with the DALI program. However, a comparison of the sequence conservation pattern of the YjeF-N domains with that of the conventional Rossmann-fold dehydrogenases reveals several notable differences (Fig. 4 and Additional file 1). These include: 1) All members of this family contain two additional consecutive N-terminal helices that precede the first strand of the β/α core of the Rossmann fold and the core itself contains eight β/α units. Both these helices contain nearly absolutely conserved acidic residues. 2) The α/β core contains two characteristic aspartates; an absolutely conserved D at the end of strand 5 and one nearly universal D at end of strand 4. 3) The first helix of the α/β core of the Rossmann fold is extended by a whole turn resulting in the abbreviation of the glycine-rich nucleotide binding loop of the fold (Fig. 4). 4) The central sheet of the Rossmann fold is highly curved to form a peculiar barrel-like structure and the second additional N-terminal helix and the first helix of the α/β core pack against each other (Fig. 4). This structural quirk is chiefly stabilized by two sets of highly conserved interactions. Firstly, the salt-bridge and hydrogen-bonding interaction between the conserved acidic residue in the second N-terminal additional helix and the RH doublet in the first helix of the α/β core helps to positioning these two helices against one side of the curved sheet. Secondly, the hydrogen bonding between the conserved aspartate at the end of strand 4 and the nearly absolutely conserved threonine C-terminal to strand 5 help in stabilizing the curvature of the central sheet (Fig. 4). 5) The acidic residue in the N-terminal-most additional helix of the YjeF-N, the acidic residue at the end of strand 5 and the polar residue (usually asparagine) from loop between strand 1 and helix 1 of the α/β core, line the mouth of the barrel-like structure to constitute the potential active site of this domain (Fig. 4).

In bacteria the YjeF-N domain is often found fused to a C-terminal kinase domain of the ribokinase superfamily (Fig. 2). Given that kinase domains are often fused to different phosphoesterase (phosphatase) domains [46], it is possible that the YjeF-N-type Rossmann fold domains may also catalyze this reaction. The conservation of the acidic residues in the predicted active site of the YjeF-N domains is reminiscent of the presence of such residues in the active sites of diverse hydrolases. Thus, in the context of the decapping pathway, it is possible that the YjeF-N domains of Dcp3p and FLJ21128 catalyze hydrolitic RNA-processing reactions, such as, phosphoester hydrolysis, dephosphorylation, demethylation or glycosyl bond hydrolysis.

The crystal structures of the archaean Sm protein, SmAP3, and MscS provide examples of Sm domain toroids with additional N-terminal and/or C-terminal domains [29,47]. These structures indicate that these extension project out on either side of the of the central heptameric toroid formed by Sm domains [29,47]. If the Scd6p were to form similar toroidal structures, then the N- and C-terminal charged extensions with RG motifs and the FDF domains of the proteins are likely to project out similarly. In the canonical Sm toroids the RNA is threaded through the central cavity of this toroid, and previous studies have suggested that the charged extensions projecting away from the Sm core may form additional non-specific contacts with the RNA [25,26,48]. A similar RNA-binding function can be envisaged for the FDF domain. However, it is also possible that it forms a distinct interaction surface to bind charged peptides from proteins belonging to a specific RNP complex, possibly the complex that is involved in decapping [45].
Scd6p and Dcp3p in the context of the origin and evolution of the decapping machinery

The provenance of the decapping-dependent RNA degradation system in eukaryotes appears to have involved a number of different innovations and recruitment events. One process involved the de novo "invention" of new \(\alpha\)-helical domains that mediate particular interactions, which are specific to this system. The most prominent of these inventions are the FDF domain and PATADs (for PAT1 alpha helical domains), the conserved \(\alpha\)-helical domains seen in yeast Pat1p and its relatives from other eukaryotes. Sequence analysis and structure prediction also suggests that the decapping proteins, Edc1p/Edc2p [53], are also potential examples of poorly structured proteins that appear to be de novo innovations of the eukaryotes. In other instances, distinctive variants of preexisting globular folds appear to have been recruited for novel functions. An example of this is the decapping enzyme subunit Dcp1p, which contains a divergent variant of the peptide-binding EVH1 domain [54] that appears to have been recruited for a different, possibly catalytic function in the decapping process.

The MutT domain of Dcp2p [55] and the YjeF-N domain of Dcp3p appear to represent cases where the ancestral active site residues of the pre-existing catalytic domains appear to have been maintained, but they acquired a new set of substrates, specific to the decapping process. Analysis of phyletic patterns shows that Dcp2p is conserved throughout currently-sampled eukaryotes suggesting that it was present in the common ancestor of the extant eukaryotes. The closest relatives of this MutT domain are seen in bacteria, suggesting that the precursor of the Dcp2p catalytic domain may have been acquired very early in eukaryotic evolution via a transfer from a bacterial lineage. The precursor of Dcp3p and FLJ21128 was probably present at least since the common ancestor of the fungi and animals. Analysis of phyletic patterns of YjeF-N domains indicates that a second version of this domain, which is not fused to the FDF domain, is conserved across the three principal superkingdoms of life. Phylogenetic analysis of this version supports the monophyly of the YjeF-N domain in each of the three superkingdoms (barring certain lateral transfer involving bacteria; data not shown), suggesting that a single copy of the YjeF-N domain is traceable to the last universal common ancestor of all life forms. Its fusion to a small-molecule kinase of the ribokinase superfamily in bacteria suggests that the ancestral form of the YjeF-N domain may have functioned in the metabolism of a critical low molecular weight compound. The version of the YjeF-N domain found in Dcp3 and FLJ21128 was probably derived in the common ancestor of the animals and the fungi through duplication of the more ancient version of the YjeF-N domain. Alternatively, it could have been acquired via lateral transfer from a bacterial lineage. The extensive sequence divergence of the two versions currently prevents us from distinguishing between these possibilities through phylogenetic analysis.

The Sm domain is an ancient RNA binding domain that appears to have bound RNA ligands even in the last uni-
versal common ancestor of all extant life forms [1,24,37,49]. In bacteria, at least two ancient versions are present, namely Hfq [50,51] and the YhbC [52] (an uncharacterized protein found in most bacteria in the same operon with genes for the translation elongation fac-
tor NusA and initiation factor IF2; VA and LA, unpublished observations). Both these versions of the Sm domain are predicted to participate in binding RNAs in the context of translation. In archaea too the Sm domains interact with various RNA ligands, such as the RNase P ribozyme [49].

The Sm superfamily of domains appears to have been vertically inherited by the eukaryotes from the common ancestor of the archaeo-eukaryotic lineage [1,37,49]. In eukaryotes the superfamily underwent a proliferation and appears to have been recruited as the core protein compo-
ment of various eukaryote-specific RNP complexes such as the spliceosomal particles, the decapping complex and the telomerase complex. Phyletic patterns suggest that their explosive diversification in eukaryotes, giving rise to highly divergent forms such as the Scd6p family, appears to have happened prior to the divergence of the extant eukaryotic lineages. This suggests that the diversification of Sm-domain superfamily might have enabled them to interact with a diverse range of RNA ligands and protein partners and there by favored the emergence of multiple eukaryote-specific RNP complexes. Subsequently each of these complexes may have developed further, through the process of innovation of new α-helical domains and recruitment of catalytic domains from various sources.

Conclusions

We show that the Scd6p family contains a novel divergent version of the RNA-binding Sm domain and a previously uncharacterized C-terminal domain, the FDF domain. While the Scd6p Sm domain is predicted to bind RNA like most other prokaryotic and eukaryotic Sm domains, it is likely to have certain unique characteristics in terms of target specificity. The FDF domain is also present in several proteins such as Dcp3p and FLJ21128, where it is combined with the YjeF-N domain, a novel version of the Rossmann fold domain, and in some cases with another divergent version of the Sm domain. Along with other atypical Sm domains, like Ataxin-2 [24], Scd6 might form alternative Sm complexes, distinct from the, classical Sm, Lsm1-7p and Lsm2-8p complexes. A variety of contextual connections from expression, protein-protein interaction and intracellular localization data, suggest that the Scd6p, Dcp3p and FLJ21128 are associated with mRNAs in the cytoplasmic substructures and possibly regulate the stability of specific messages via the decapping system. The FDF domain may mediate interactions that are specific to these RNP complexes. Phyletic analysis of other components of the decapping system suggests that they have diverse ori-
gins and the explosive diversification of the Sm domains at the base of the eukaryotic radiation may have played an important role in the provenance of the uniquely eukary-
otic RNP complexes.

Methods

The non-redundant (NR) database of protein sequences (National Center for Biotechnology Information, NIH, Bethesda) was searched using the BLASTP program [56]. Iterative database searches were conducted using the PSIBLAST program with either a single sequence or an align-
ment used as the query, with the PSSM inclusion expecta-
tion (E) value threshold of 0.01 (unless specified otherwise); the searches were iterated until convergence [56,57]. For all searches with compositionally biased pro-
teins, the statistical correction for this bias was employed. Multiple alignments were constructed using the T_Coffee [58] or PCMA [59] programs, followed by manual correc-
tion based on the PSI-BLAST results. Globular domains were predicted using the SEG program with the following parameters: window size 40, trigger complexity = 3.4; extension complexity = 3.75 [22]. All large-scale sequence analysis procedures were carried out using the SEALS package [60]. Specifically, pattern searches were carried out using the gref program from this package. Structural similarity searches were conducted using the DALI pro-
gram. The Swiss-PDB viewer [61] and Pymol programs were used to carry out manipulations of PDB files. Figures were rendered using PyMOL [62,63] or PoV-Ray [64].

Protein secondary structure was predicted using a multi-
ple alignment as the input for the PHD program [65,66]. Similarity-based clustering of proteins was carried out using the BLASTCLUST program [67].

Phylogenetic analysis was carried out using the maxi-
mum-likelihood methods. Maximum-likelihood distance matrices were constructed with the TreePuzzle 5 program [68] using 1000 replicates generated from the input align-
ment and used as the input for construction of neighbor-
joining trees with the neighbor program [69]. Neighbor uses a weighted N tree construction procedure that has been shown to effectively correct for long-branch effects [69]. Alternatively a full ML tree was constructed using the Proml program of the Phylip package [70]. This tree was used as the input tree to generate further full ML trees using the PhyML program [71] with 100 bootstrap replicates generated from the input alignment. The consensus of these trees was derived using the Consense program of the Phylip package to obtain the bootstrapped ML tree. Gene neighborhoods were determined by searching the NCBI PTT tables with a custom-written script. These tables can be accessed from the genomes division of the Entrez retrieval system [72].
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