Multicomponent 2′-O-Ribose Methylation Machines: Evolving Box C/D RNP Structure and Function
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Abstract

Methylation at the 2′-O ribose position is an abundant nucleotide modification of both eukaryal and archaeal RNAs. The methyltransferase responsible for this modification is frequently a ribonucleoprotein (RNP) complex consisting of a box C/D guide RNA and associated core proteins. These RNP “machines” are responsible for the modification of numerous cellular RNAs including ribosomal RNA, spliceosomal snRNAs and transfer RNAs. This chapter will review the structure and function of both eukaryotic and archaeal box C/D RNPs. A particular focus of our discussion will be the evolving components of the box C/D RNPs and the resultant consequences upon box C/D RNP structure and function.

Introduction

Guide RNAs for nucleotide modification were first described in the eukaryotic nucleolus where they were shown to modify ribosomal RNA. Based upon conserved sequence elements, these small nucleolar RNAs (snoRNAs) were classified into two major families. The box C/D snoRNAs guide nucleotide 2′-O-methylation whereas the H/ACA snoRNAs isomerize uridine to pseudouridine. Subsequent investigations revealed that box C/D and H/ACA guide RNAs are also found in Archaea. Further characterization of both eukaryotic and archaeal guide RNAs has demonstrated that they are bound by core proteins to form ribonucleoprotein (RNP) complexes. Both RNP families accomplish nucleotide modification using a similar mechanism. Guide RNAs utilize complementary sequences to base pair with specific target RNAs, thus designating a specific nucleotide for modification. The RNA-bound core proteins catalyze the 2′-O-methyl transfer and pseudouridylation reactions. The focus of this chapter is the evolving structure and function of the box C/D RNPs. For a detailed discussion of the H/ACA RNP structure and function, the reader is referred to Chapter 22 by Grozdanov and Meier entitled “Multicomponent Machines in RNA Modification: the H/ACA Ribonucleoproteins”.

Ribonucleotide Methylation and Methylation Function

Key features of ribose 2′-O-methylation indicate that this abundant nucleotide modification plays an important role in RNA folding and stability. Methylation at the ribose 2′ position stabilizes an RNA chain by inhibiting backbone cleavage and increasing the stability of base pairing and stacking interactions, thus potentially affecting the RNA's structure and ultimately function. A number of important cellular RNAs are 2′-O-methylated by box C/D RNPs. Although the

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function of tRNA modification is not fully understood, disrupting box C/D snoRNA-directed 2′-O-methylation results in slowed cell growth and reduced ability of cells to adapt to environmental changes with adverse affects on ribosome biogenesis and function.3,6 When mapped on the ribosome, these modifications cluster around functionally significant regions like the peptidyl transferase center.4 Eukaryotic spliceosomal RNAs (snRNA) are also methylated by a variety of box C/D snoRNAs as are small Cajal RNAs (scaRNAs), RNAs localized to nuclear Cajal bodies that can contain both box C/D and H/ACA motifs.7 Box C/D snoRNA-directed methylation of select mRNAs has been implicated in regulating RNA editing and splicing of brain mRNAs.8,9 Computational analyses have recently revealed that alternative splice junctions may also be targets for snoRNA-guided modification as they are often complementary to a number of “orphan” box C/D snoRNA guide regions.10 Unique to Archaea, box C/D RNAs guide methylation of tRNAs, thus potentially affecting not only tRNA folding and structure but also tRNA function in translation.11,12

Box C/D RNP Function
The primary function of eukaryotic and archaeal box C/D RNPs is nucleotide methylation of diverse cellular RNAs. However, other functions in RNA metabolism have been demonstrated. In eukaryotes, box C/D snoRNAs function in prerRNA processing. Select box C/D snoRNAs are essential for specific endonucleolytic cleavage events in prerRNA maturation, likely functioning as "organizers" for a trans-acting RNase.13-15 Several box C/D snoRNAs also play roles in prerRNA folding.15-17 For both functions, the box C/D snoRNA utilizes complementary sequences to base pair with the prerRNA. Notably, these additional functions have not yet been observed for archaeal box C/D snoRNAs.10 This may reflect a more limited examination of archaeal box C/D sRNA populations and functions or perhaps evolving functional roles of the box C/D snoRNP in eukaryotes.

Box C/D RNAs: Diversity of Sequence and Structure
Large populations of box C/D RNAs are found in eukaryotic and archaeal organisms. In various archaeal organisms, scores of box C/D sRNAs have been identified using bioinformatic approaches and many have been experimentally verified.19-22 However, the list of sRNAs remains small and is still limited to a handful of organisms. It appears that Archaea do not share box C/D RNA homologs with Eukarya, indicating an evolutionary ancient divergence of eukaryotic and archaeal RNAs.19,23,24 Box C/D snoRNA populations are better defined in eukaryotes, although not nearly complete. In the unicellular eukaryote yeast, the defined box C/D snoRNA population consists of 46 species.25-26 In humans, a larger population of over 100 box C/D snoRNAs has been identified and this number is likely to grow significantly.24 Interestingly, the identification of brain-specific species in mammals suggests an expanding complexity of tissue-specific RNAs and perhaps snoRNA function in metazoan organisms.9,27,28 Even more numerous may be the plant box C/D RNAs whose populations are predicted to be in the hundreds.29 Although box C/D snoRNAs from different eukaryotic organisms can guide evolutionarily conserved modifications, most nucleotides targeted for modification are unique to a given organism, reflecting the general lack of snoRNA species conservation among eukaryotes.

The hallmark of box C/D guide RNAs are the box C (RUGAUGA) and box D (CUGA) sequence elements located at the 5′ and 3′ RNA termini, respectively (Fig. 1A). Frequently present are internal box C′ and D′ elements which are well conserved in archaeal sRNAs but often difficult to discern in eukaryotic snoRNAs.30 These terminal and internal boxes establish the box C/D and C′/D′ motifs, respectively. Both motifs fold into RNA elements known as kink-turns (K-turns) first revealed in U4 snRNA and archaeal ribosomes.31,32 K-turns are characterized by an asymmetric bulge flanked by two stems and stabilized by tandem, sheared G:A pairs. The G:A pairs hydrogen bond across the bulge to generate a sharp, archetypical bend, or kink of approximately 60° in the RNA backbone.31 Importantly, internal C′/D′ motifs fold into a modified K-turn structure where canonical stem I is replaced by a loop. These modified K-turns have been
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...K-turns have also been observed in mRNAs, archaeal H/ACA sRNAs and even the SAM riboswitch. K-turn and K-loop motifs are typically protein binding platforms, important for stabilizing tertiary RNA and RNP structures. Individual box C/D RNA species are defined by their unique guide sequences located upstream of boxes D and D'. Guide sequences are...
10-21 nucleotides long and complementary to their respective target RNA. It is the target RNA nucleotide which is base paired to the fifth nucleotide of the guide sequence that is specifically 2′-O-methylated by the RNP complex.37

Differences in size and structure between the archaeal and eukaryotic box C/D RNAs has contributed to structural and perhaps functional diversity. Archaeal sRNAs are smaller (50-70 nucleotides) and possess terminal box C/D and internal C’/D’ motifs separated by minimal guide regions. Guide region length is highly conserved at 12 nucleotides in archaeal box C/D sRNAs and thus box C/D and C’/D’ motif spacing is conserved.20,38 Interestingly, circular box C/D sRNAs have been reported in some archaeal organisms.39 In contrast, eukaryotic snoRNAs are larger in size (most often greater than 75 nucleotides) with significantly larger guide regions and associated spacer sequences between the two motifs. For those box C/D snoRNAs with hard to define or missing C’/D’ motifs, the D guide sequence and associated spacer region can be quite large.24,25,29,38 Some eukaryotic box C/D snoRNAs utilize their guide regions for prerRNA processing steps. The larger RNA size and correspondingly larger guide regions may have contributed to and even facilitated the functional diversity of box C/D snoRNP in Eukarya.

Box C/D RNP Structure and Assembly

Mature box C/D RNAs are assembled as ribonucleoprotein complexes bound with a limited number of highly conserved core proteins (Fig. 1B). Eukaryotic box C/D snoRNPs contain four conserved core proteins: the 15.5kD protein, nucleolar proteins Nop56 and Nop58 and the methyltransferase enzyme fibrillarin.40-42 Three highly homologous proteins, ribosomal protein L7Ac, Nop56/58 and fibrillarin, bind the archaeal box C/D sRNAs to assemble a simpler and what could be considered minimal box C/D sRNP complex.43,44

In vitro reconstitution of catalytically active archaeal box C/D sRNPs has revealed an order of core protein binding.44,45 L7Ac initiates sRNP assembly by binding the K-turn and K-loop motifs of the terminal box C/D and internal C’/D’ motifs, respectively.43-45 Nop56/58 and then fibrillarin bind both the terminal box C/D and internal C’/D’ motifs to assemble a “symmetric” sRNP with all three core proteins bound to both motifs.44,45 The assembly of a symmetric RNP is essential for efficient nucleotide methylation.45,46 Initial binding of L7Ac core protein stabilizes K-turn and K-loop structure and remodels the box C/D RNA to facilitate subsequent binding of the Nop56/58 and fibrillarin proteins.47,48 Remodeling of the sRNA continues with binding of Nop56/58 while fibrillarin has no significant affect on RNA structure.48 For the archaeal complex, RNA remodeling requires elevated temperature to increase RNA structure dynamics, thus facilitating core protein binding. Notably, in vitro assembly of archaeal box C/D sRNPs does not require accessory proteins for either RNA remodeling or hierarchical core protein binding.44,45

In contrast to the symmetric archaeal sRNP, the eukaryotic box C/D snoRNPs assemble an apparently “asymmetric” complex.49 The 15.5kD protein initiates snoRNP assembly but appears to bind only the K-turn of the terminal box C/D motif.50 Core proteins Nop58 and Nop56 have been predicted to bind the C/D and C’/D’ motifs, respectively, based upon in vivo crosslinking experiments.49 Only fibrillarin appears to be associated with both motifs. Unfortunately, the lack of a functional in vitro assembly system for the eukaryotic complex has hampered a more detailed analysis of box C/D snoRNP assembly and structure.

Limited knowledge of eukaryotic box C/D snoRNP assembly has nonetheless revealed a highly complex and dynamic process requiring accessory factors. Assembly of the mammalian presnoRNP requires two trans acting AAA+ ATPases, TIP48 and TIP49.51,52 Additional processing/assembly factors for the U3 snoRNP include TGS1, La, LSm proteins and the exosome as well as nucleocytoplasmic transport factors such as PHAX, CRM1, CBC, Ran and Nopp140.51,52 Four novel human biogenesis factors (BCD1, NOP17, NUFIP and TAF9), which are likely to be involved in the formation of the U8 presnoRNP, have also been identified.49 Most recently, the heat shock protein Hsp90 has been implicated in orchestrating assembly of the eukaryotic complex.55,56 Whereas archaeal sRNPs require elevated temperature (accessory factors in vivo?) to facilitate RNA remodeling required for in vitro sRNP assembly, the eukaryotic snoRNPs require multiple...
accessory factors for in vitro and in vivo assembly. These accessory factors are presumed to promote RNA remodeling and facilitate sequential core protein binding, an apparently common theme of both archaeal and eukaryotic box C/D RNP biogenesis.

The vast majority of higher eukaryotic snoRNA coding sequences are positioned within introns of RNA Polymerase II protein coding host genes. A second genomic organization, prevalent in yeast and plants, is box C/D snoRNA genes transcribed from independent RNA Pol II (infrequently Pol III) promoters. 57 Archaeal box C/D sRNA genes, although not well characterized, appear to be intergenic and transcribed from independent promoters. 51,20 Transcription of intronic box C/D snoRNAs is coupled with the transcription of the host premRNA and linked to splicing. 58 Box C/D snoRNP assembly is also coupled with posttranscriptional processing, maturation and transport to the nucleolus. 59 The differences in genomic organization for the eukaryotic box C/D snoRNAs versus archaeal sRNA coding sequences perhaps reflects an evolution of gene structure for purposes of regulated expression.

Structure, Function and Evolution of the L7Ae/15.5kD Core Protein

Archaeal core protein L7Ae initiates sRNP assembly by binding the terminal box C/D (K-turn) and internal C'/D' (K-loop) motifs. 44,45 L7Ae binding remodels sRNA structure and establishes a platform for subsequent box C/D sRNP core protein binding. 47,48,60 Eukaryotic 15.5kD protein similarly initiates snoRNP assembly by binding the terminal box C/D core motif’s K-turn. 33,45,50 The differential binding of L7Ae and 15.5kD proteins to K-turn and K-loop motifs in vitro is striking as the crystal structures of both proteins are nearly superimposable and their RNA-binding domains are well conserved across both domains of life. 31,61,62 (Fig. 2).

L7Ae and 15.5kD are members of the L7Ae/L30 protein family. 31,61,62 (Fig. 2). Additional members of this family include rpL30e in Archaea and Rpp38, rpL30, rpl7a, SBP2 and Nhp2p proteins in Eukarya. Proteins in this closely related family are typically small and composed of an internal beta sheet surrounded by several alpha helices, a three-layer topology fold known as an alpha-beta-alpha sandwich. 32,34 They possess conserved RNA binding domains, almost uniformly recognize K-turn motifs and play critical roles in RNA stabilization and RNP assembly. 31,32,34,42,55,61-63

Each family member is interesting from a functional standpoint. Family members in both Archaea and Eukarya function as ribosomal proteins of the large subunit. Eukaryotic ribosomal protein rpL30 is also capable of binding its own mRNA to regulate translation and ribosomal protein Rpp38 is a constituent protein of the MRP complex. 34,64 SBP2 is another mRNA-binding protein, recognizing those mRNAs possessing the SECIS RNA element important for selenocysteine incorporation into selenoproteins. It consists of multiple domains including one very similar to that found in the L7Ae/15.5kD protein. 65 An L7Ae/L30 sequence appears to have been inserted during genomic shuffling, thus conferring K-turn RNA binding capability upon SBP2. 35 The 15.5kD protein is not only a box C/D snoRNP core protein but also a component of the spliceosomal U4 snRNP where it also binds a K-turn motif in U4 and functions in snRNP assembly. 32,42,60 Eukaryotic nonhistone chromosomal protein 2 (NHP2p) is a core protein of the eukaryotic H/ACA snoRNPs and highly homologous to both archaeal L7Ae and eukaryotic 15.5kD proteins. NHP2p binds a stem loop of the box H/ACA snoRNAs and is essential for H/ACA snoRNP assembly. Notably, NHP2p stands out as being the sole L7Ae/L30 family member without clear RNA-binding specificity. Specific recruitment of NHP2 to the assembling snoRNP requires interaction with the RNA and other core proteins. 66 Its functional equivalent in the archaeal H/ACA sRNP is L7Ae, the only guide RNA core protein of both domains to be found in both the box C/D and H/ACA RNPs and binding the K-turn/K-loop motifs. Despite great similarities in sequence and folded structure, each L7Ae/L30 family member has sufficiently diverged such that its binding is specific for the K-turn of its respective cognate RNA. 31,34,35,45,50,66

The recurring theme of L7Ae/L30 protein function is RNP formation via recognition of the K-turn motif. The binding of L7Ae/L30 proteins to a variety of RNAs provides insight into the evolutionary emergence of the L7Ae/L30 protein family and even evolution of the box C/D RNP s. The limited number of L7Ae/L30 proteins in Archaea (two) and expansion of family members in
eukaryotes (six) suggest a continuing evolution and diversity of protein structure and function, particularly in eukaryotic organisms. L7Ae is a component of three separate RNPs in Archaea whereas in eukaryotes these same functions are carried out by three separate but closely related family members (ribosomal protein L7a, 15.5kD, NHP2). This would suggest that L7Ae is the progenitor of the L7Ae/L30 protein family.31,43,45,50,62 We have previously proposed that L7Ae or an L7Ae‑like protein binding a K‑turn motif in a primitive RNP translational apparatus may be the ancestral RNP complex for this protein family.23 The utilization of a single archaeal core protein to bind K‑turns in both archaeal box C/D and H/ACA RNPs suggests a common RNP origin for both guide RNP families early in evolution. The absence of L7Ae/L30 RNP complexes in Eubacteria implies an emergence and evolution of the protein family after divergence of Archaea and Eukarya from Eubacteria.
Evolution of L7Ae/15.5kD RNA-binding capabilities may well have facilitated evolution of box C/D RNP structure and hence function. The minimal archaeal box C/D sRNP makes use of L7Ae for both K-turn and K-loop binding, thus assembling a symmetric sRNP whose box C/D and C'/D' RNPs are spatially constrained and functionally coupled. In contrast, the evolved binding capability of eukaryotic 15.5kD to recognize only the box C/D K-turn could have allowed greater structural and consequently functional snoRNP diversity. Lack of sequence conservation in the C'/D' motif of eukaryotic snoRNAs may reflect a concomitant loss of 15.5kD binding, resulting in spatial decoupling of the internal and terminal motifs. Thus, modern day eukaryotic box C/D snoRNAs are less conserved in sequence and larger in size. This flexibility in snoRNA structure may have allowed the eukaryotic complexes to drift further, acquiring new functions such as chaperoning preRNA processing events.

Structure, Function and Evolution of the NOP56 and NOP58 Core Proteins

The Nop core proteins play essential structural and functional support roles in the box C/D RNPs. A single Nop56/58 is found in Archaea while two homologs, presumably arising from gene duplication and designated Nop56 and Nop58, are present in Eukarya. Their roles include bridging protein interactions within box C/D RNPs, RNA remodeling during RNP assembly, fibrillarin recruitment and assisting the methyltransferase reaction. Archaeal Nop56/58 helps to remodel RNA structure during in vitro box C/D sRNP assembly by restructuring guide regions and box elements after initial remodeling by L7Ae. Nop56/58 interactions with other core proteins may affect RNA remodeling, perhaps helping to establish bridging interactions between the box C/D and C'/D' RNPs. Archaeal Nop56/58 and eukaryotic Nop56 and Nop58 proteins interact with fibrillarin. Evidence from in vitro assembly of the archaeal sRNP suggests that Nop56/58 and fibrillarin may bind the assembling complex as a dimer. While the methyltransferase fibrillarin clearly interacts with guide and target RNAs, its binding in the archaeal sRNP is primarily through interaction with box C/D RNPs. Archaeal Nop56/58 may also assist in catalysis of the methyltransferase reaction as critical Nop56/58 amino acids are positioned adjacent to the S-adenosyl-L-methionine binding site of fibrillarin.

Only a few members of the Nop protein superfamily have been well characterized. They include the box C/D RNP core proteins Nop56/58 in Archaea, Nop56 and Nop58 in Eukarya and eukaryotic Prp31 (premRNA processing factor 31). Nop proteins are composed of an N-terminal domain, a central coiled-coiled domain, a Nop domain and a variable lysine-rich C-terminal tail (Fig. 3A). The N-terminal domain is not well characterized in eukaryotes but is responsible for dimerization with fibrillarin in Archaea. The coiled-coil domain may mediate protein interactions with other core proteins or regulatory factors. Crystal structures of the Nop56/58-fibrillarin dimer from Archaea show that the coiled-coil domain can dimerize with itself, leading to the suggestion that this interaction could mediate protein-protein or crosstalk interactions between the box C/D and C'/D' RNPs. Best understood is the Nop domain, the defining feature of the Nop superfamily, which comprises most of the C-terminal region. A recent U4-15.5kD-Prp31 RNP crystal structure has provided new insight into the role of this domain in RNP assembly. (Fig. 3B). The Prp31 Nop domain makes nearly equal contact with both U4 RNA and the 15.5kD protein, thus explaining a need for 15.5kD to be bound to U4 for Prp31 interaction. In a similar manner, archaeal Nop56/58 binds a box C/D RNA only after L7Ae has first bound the K-turn or K-loop motif. Thus, the Nop proteins may serve as checkpoints in RNP assembly, ensuring that the K-turn recognition protein has first bound RNA. Deletion of the Nop domain completely disrupts binding to the box C/D RNA-L7Ae complex, indicating that it is the necessary RNP assembly module of Nop protein family members. The highly charged, lysine-rich C-terminal tail, also called a KKE/D repeat, remains an enigma. It is poorly conserved in sequence and length and appears to be dispensable for Nop protein function in both Eukarya and Archaea.
Eukaryotic box C/D snoRNPs may owe much of their structural and functional diversity to evolution of the Nop56/58 core protein. In archael box C/D sRNPs the Nop56/58 protein binds both box C/D and C'/D' motifs. In contrast, crosslinking experiments indicate that eukaryotic Nop56 and Nop58 may differentially bind the C'/D' and C/D motifs, respectively. Nop56 and Nop58 are highly related, with the mouse proteins having 43% identity and 63% similarity. Archaeal Nop56/58 from Methanocaldococcus jannaschii is 57% and 59% similar to mouse Nop56 and Nop58, respectively. Thus, gene duplication of Nop56/58 coding sequence followed by co-evolution of the two eukaryotic proteins and the box C/D RNA could contribute to the apparent asymmetric structure of eukaryotic box C/D snoRNPs. As 15.5kD does not recognize the K-loop, association of Nop56 with the C'/D' motif could suggest that this Nop protein has acquired the ability to bind RNA independently of 15.5kD. In vitro assembly of the archael sRNP has also shown that archael Nop56/58 along with fibrillarin can specifically, albeit weakly, bind the K-loop motif in the absence of L7Ae RNA. The possible differential recognition of Nop56 and Nop58 proteins to K-loop and K-turn motifs, respectively, as well as the K-turn specificity of the 15.5kD protein, could also contribute to the uncoupling of the eukaryotic box C/D and C'/D' RNP complexes.
Structure, Function and Evolution of Fibrillarin

Fibrillarin is the catalytic protein of the box C/D RNPs, yet it plays only a minor role in RNP assembly. In Archaea, fibrillarin is recruited to the complex primarily through protein-protein interaction with the Nop56/58 protein. In eukaryotes, fibrillarin may play a more active role in assembly. Eukaryotic fibrillarin contacts the box C/D snoRNAs and association of Nop56 requires the presence of fibrillarin. Fibrillarin is recruited to the RNP at a late stage of assembly.

Fibrillarin was originally predicted to be the methyltransferase enzyme based on its sequence similarity to other S-adenosyl-l-methionine (SAM)-dependent methylases. Subsequent in vitro reconstitution of box C/D RNPs and crystallographic analyses of archaeal fibrillarins provided further evidence of the methyltransferase function of fibrillarin. Despite this progress, it is still unknown exactly how fibrillarin interacts with guide and target RNAs to accurately methylate the target nucleotide.

Eukaryotic and archaeal fibrillarin proteins have both common and unique features. They all share a highly conserved alpha-beta carboxy-terminal domain (CTD) in which is nested a short consensus sequence, the SAM-binding motif. The CTD of M. jannaschii fibrillarin (Mjfib) is approximately 60% identical and 80% similar to vertebrate fibrillarins between residues 25 and 95 of the CTD, which harbors the SAM-binding motif. Even in poorly related regions outside this segment (Mjfib residues 95-227), archaeal and eukaryotic fibrillarins are about 40% identical and 65% similar.

In contrast to the CTD, fibrillarin proteins have variable sequence and structure in their N-terminal domains (NTD). Eukaryotic fibrillarins often contain a glycine-arginine-rich (GAR) domain which is necessary and sufficient for nucleolar localization of eukaryotic box C/D snoRNPs. However, archaeal fibrillarins lack this domain and their N terminal regions are much shorter (Fig. 4A). Moreover, the fibrillarin NTD varies within archaeal species and may confer different protein binding properties upon them. For example, the Mjfib NTD was reported to facilitate dimerization of fibrillarin molecules through specific β-strand interactions. In contrast, available evidence indicates that fibrillarin from both Archaeoglobus fulgidus and Pyrococcus furiosus exist as monomers in solution and in crystalline state.

Despite a lack of significant sequence homology, the archaeal fibrillarin CTD is structurally similar to other SAM-dependent methylases. The consensus topology for the methyltransferase catalytic domain is a seven-stranded β-sheet flanked by three α-helices on each side. The CTD of MjFib forms a Rossman fold like other methyltransferases and only differs from the consensus topology by the addition of a minihelix (α5). Fibrillarin is most closely related to other SAM-dependent RNA methyltransferases, like RrmJ from E. coli which catalyzes site-specific 2′-O-methylation of rRNAs, tRNAs and mRNAs independent of a guide RNA. The site-specific RNA methyltransferases (MTases) related to RrmJ and snoRNA-directed RNA MTases related to fibrillarin form a closely related monophyletic clade. They possess a spatially superimposable tetrad of conserved residues localized in the heart of the substrate-binding pocket, three of which (K-D-K) are essential for activity. This invariant triad is considered a synapomorphy, an ancient feature derived from a common ancestor that might have possessed ribose 2′-O-MTase activity. Collectively, these observations suggest that methyltransferase enzymes evolved from a common ancestor to acquire substrate-specific activities.

Fibrillarin relies upon a guide RNA and other core proteins in an assembled box C/D RNP to catalyze nucleotide-specific 2′-O-methylation. Most other methyltransferases utilize accessory domains for substrate specificity. For example, the DNA methylase HhaI recognizes and binds its double-stranded DNA substrate by utilizing a large peripheral domain which binds the DNA and flips the target base out of the duplex for modification. Evolution of fibrillarin appears to have occurred within the box C/D RNPs as well. Archaeal fibrillarins possess organism-specific NTDs while eukaryotic fibrillarins have related GAR domains. Aside from affecting nucleolar localization, the GAR domain serves as an interaction domain with the SMN protein which is transiently associated with premature box C/D snoRNPs and important for assembly.
Interestingly, eukaryotic fibrillarin may have other roles in addition to ribose methylation. Most eukaryotic box C/D snoRNPs appear to direct only one ribose methylation per snoRNA.

Figure 4. Conserved sequence and structure of fibrillarin. A) Sequence alignment of three eukaryotic and three archaeal fibrillarins with the *E. coli* RrmJ methyltransferase. Degree of conservation is indicated by shades of gray. The highly conserved SAM-binding motif is boxed. B) Crystal structure of *M. jannaschii* fibrillarin (1FBN). The variable N-terminal domain is light gray, the SAM-binding motif circled and highly conserved catalytic residues designated (black sticks). C) Spatial superposition of the *E. coli* RrmJ catalytic residues (black) (1EIZ) with those of *M. jannaschii* fibrillarin (light gray). The invariant catalytic triad (K-D-K) is labeled and peptide backbones are illustrated with lines.

Interestingly, eukaryotic fibrillarin may have other roles in addition to ribose methylation. Most eukaryotic box C/D snoRNPs appear to direct only one ribose methylation per snoRNA.
even though fibrillarin is believed to bind both box C/D and C’/D’ motifs. Notably, box C/D snoRNPs involved only in prerRNA processing or folding, such as U3 and U8, also contain the fibrillarin core protein.54,83 These observations suggest that eukaryotic fibrillarin may have acquired a more structural role in some RNPs and may possess other functions aside from strictly catalyzing the methyltransfer reaction.

Concluding Remarks: The Evolving Box C/D RNP Machinery

RNA-guided nucleotide modification complexes are ancient RNA:protein enzymes found in both Eukarya and Archaea. Despite their conservation in these two domains of life, the box C/D RNPs exhibit domain-specific structural and functional features indicating an evolving RNP over time. The archaeal sRNP complex can well be considered a minimal RNP composed of smaller RNAs, three core proteins, with spatially and functionally coupled box C/D and C’/D’ RNPs. Known RNA targets are confined to ribosomal and transfer RNAs and its only function appears to be nucleotide modification. The sRNAs are directly transcribed from intergenic genes and assembly of the sRNP does not require, at least in vitro, accessory proteins. In contrast, the eukaryotic snoRNP is more complex both structurally and functionally. It is composed of larger RNAs, one additional core protein resulting from gene duplication, with poorly conserved C’/D’ RNPs that do not appear to be spatially linked to the box C/D RNP. SnoRNP target RNAs are more diverse and RNP functions include rRNA folding and processing as well as nucleotide modification. The snoRNA genes are varied in genomic organization, often transcribed as introns and snoRNA processing is essential with RNP assembly requiring numerous assembly factors.

In this chapter, we have presented the current state of knowledge concerning the structure and function of the box C/D RNPs. Our focus has been comparison of the archaeal and eukaryotic complexes, detailing their differences to provide the reader with an overview of the evolving box C/D RNP complexes. More remains to be learned about these ancient enzymes and the coming years are certain to yield exciting and unexpected findings.

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