Distribution, organization an evolutionary history of La and LARPs in eukaryotes

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ABSTRACT
The fate of any cellular RNA is largely influenced by the nature and diversity of its interactions with various RNA-binding proteins (RBPs) leading to the formation of a biologically significant ribonucleoprotein (RNP) complex. La motif-containing proteins (composed of genuine La and La-related proteins (LARPs)) represent an evolutionary conserved family of RBPs that encompass a large range of crucial functions, involving coding and non-coding RNAs. In this work, we provide data that extend our previous knowledge on the distribution, organization and evolutionary history of this important protein family. Using a repertoire of 345 La motif-containing proteins from 135 species representing all major eukaryotic lineages, we were able to pinpoint many lineage-specific variations in the structural organization of La and LARPs and propose new evolutive scenarios to explain their modern genomic distribution.

Introduction

RNA-binding proteins (RBPs) are key determinants of coding and non-coding RNA metabolism. La motif-containing proteins are part of an important RBP family that emerged early in eukaryotic evolution and acquired specialized functions [1–3]. The genuine La protein (also named the La autoantigen) is the most ubiquitous member of this family. Genuine La proteins have been shown to protect the 3′-end of nascent RNA polymerase III transcripts and, in many occasions, to act as an RNA chaperone to prevent RNA misfolding [4,5]. La-Related Protein (LARP) 7 targets specialized non-coding RNAs (Vertebrates 7SK [6,7] or Ciliates [8,9] and Fungi [10–12]) to promote their assembly into a functional ribonucleoprotein complex. Other LARPs (1, 4 and 6), as well as the cytoplasmic isoform of the genuine La, target mRNAs and regulate their cellular localization, stability and translation potential [13–19]. Several extensive recent reviews have been published on the functions of genuine La and LARPs [1,13,20] and this aspect will not be developed further here. In light of the recent accumulation of genomic data, the objective of this work is to re-evaluate the distribution of genuine La and LARPs using data from 135 species, representing all major eukaryotic lineages (Metazoa, Embryophytes, Chlorophytes, Fungi, Alveolates, Excavates, Stramenopiles, Rhizaria and Hacrobia). We also determined the phylogenetic relationships of the 345 La and LARP proteins collectively and revisited the structural organization of each subfamily.

Results

Genuine La

A ‘classical’ genuine La protein contains a La motif, closely followed by a first RRM (RRM1), by a more distal second atypical RRM (RRM2) and by a C-terminal GK/R-rich basic motif. The combination of the La/RRM1 is usually referred to as the La Module (LaM). This classical organization is found in all Chlorophytes, Embryophytes and animal (vertebrate and invertebrate) species without any detected exception. All animal genomes possess a single genuine La gene. However, in Embryophytes, although Bryophytes, Lycophytes, basal vascular plants and most Eudicots present a single La gene per species, all Poaceae (BstLa, OsLa, PhLa, SiLa) and Brassicaceae (AhLa, AtLa, BSla, ESla) species have two (Figure 1). In Fungi, the organization of genuine La proteins can differ from the classical one described above. For example, Ascomycota La are usually shorter and do not possess an RRM2. However, other Fungi possess longer genuine La, many of which present a conserved RRM2. This is the case for many Basidiomycota (CcLa, LbiLa, MosLa, PgLa, WhLa, XdLa), Mucoromycota (EspLa, LtrLa, RiLa) and Zopagomycota (BmeLa) genuine La proteins that have a ‘classical’ organization (Figure 1). The situation in basal eukaryote lineages is also variable. While genuine La proteins from Stramenopiles (PsLa, TcLa), Amoebozoa (HaLa, DdLa), Rhizaria (PbLa) and Excavates (NgLa) have the classical structure (LaM, RRM2, GK/R-rich motif), Hacrobia (EbLa, GtLa) La miss the RRM2 (Figure 1). The organization of what appears to be Alveolates genuine La proteins is quite unusual. These proteins (PmLa, SleLa, ThLa, TgLa) contain a La motif, clearly related to other eukaryotic genuine La proteins, but no RRM nor GK/R-rich motif. Instead, they possess a conserved C-terminal region of unknown function named DUF3223 (see Figure S1). The La/DUF3223 association is specific to Alveolates, with a single-identified exceptio for the Hacrobia species Chrysochromulina tobinii that also possesses a La/DUF3223 containing protein (Figure S1)

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LARP1

Genes encoding LARP1 were detected in Amoebozoa, Fungi, Embryophytes, Chlorophytes and Metazoans. Since genomic data from several basal eukaryotic lineages are still scarce, the presence of LARP1 in other eukaryote lineages than the ones reported here cannot be completely excluded at this time. LaM, DM15, as well as two additional regions (named C-terminal Region 1 and 2 (CR1 and CR2)), are conserved in all full-length eukaryotic LARP1s. The RRM region, adjacent to the La motif, is conserved at the primary sequence level only in closely related vertebrate species. In other eukaryotes, its presence was inferred solely from secondary structure predictions. The putative unconventional PAM2 motif, identified as part of the human LARP1 RRM region, is only conserved in vertebrates (see Figure S2). In Embryophytes and Fungi, in addition to full-length LARP1, shorter LARP1 versions, with a C-terminal La motif as the sole conserved region, can also be found. While basal Embryophyte (AtrLARP1, MpoLARP1, PhpLARP1), as well as Chlorophyte (CrLARP1) species have only one full-length LARP1, all Angiosperms possess one full-length and one to several of these solo La motif versions of LARP1. In Fungi, the situation is more complex. In basal Fungi lineages (Cryptomycota (PsLARP1), Zoopagomycota (BmeLARP1, SmELARP1, TsPLARP1) and Mucoromycota (ArLARP1, LtrLARP1, RiLARP1)), only full-length LARP1 is present while in Basidiomycota, only shorter versions (with a C-terminal solo La motif) are present. In Ascomycota, the situation is intermediate, with two sub-lineages (Saccharomycotina and Taphrinomycotina) having only shorter versions (with the curious exception of one Taphrinomycotina species Neoleata irregularis (NiLARP1) that has only a full-length LARP1), and one sub-lineage (Pezizomycotina, DfLARP1, EgrLARP1, GlLARP1, McLARP1, UpLARP1) having both the full-length and shorter types. However, full-length LARP1 from Pezizomycotina are less conserved compared to their orthologues from basal Fungi lineages (illustrated by long branches for these species in Figure 2). Other LARP1 lineage-specific variations include a very short N-terminal region for Amoebozoa LARP1s (DdLARP1, HaLARP1, TILARP1) and the presence of conserved glycine-arginine (GR) repeats, N-terminal to the La motif, for Metazoans and (full-length) Fungi LARP1. A gene duplication event also likely occurred in the common ancestor of vertebrates generating two distinct versions of full-length LARP1 (named LARP1A and B, grey shading in Figure 2). This is the only situation where two distinct full-length LARP1 versions co-exist in the same lineage.

LARP4

Genes coding for LARP4 are found in Amoebozoa, Stramenopiles, Excavates as well as Metazoans, but are apparently absent from other eukaryotic lineages, including Embryophytes, Chlorophytes and Fungi. Most vertebrates...
possess two related but distinct LARP4 versions (LARP4A and B) while some fishes (but not all) have three versions (two of the A types (A1 and A2) and one of the B types) (see Figure 3). The LARP4 LaM module is the only region conserved in all LARP4s, while Metazoans present one to several other conserved motifs. Indeed, invertebrate LARP4s have a single supplementary conserved region (CR1, Figure S3), while vertebrate LARP4s have four (PAM2 w, PMB, CR1 and CR2). The existence of a consensus PAM2 motif in some invertebrates was observed\(^{[22]}\), but overall this motif is not present in invertebrate LARP4s. Human LARP4A PAM2 w and PMB have been shown to be functional PolyA-Binding Protein (PABP)-interacting regions\(^{[23]}\). Also, a receptor for activated C kinase (RACK1) interacting region (RIR) was identified near the C-terminus of human LARP4A\(^{[1]}\). It remains to be seen if this RIR corresponds to the conserved CR2 identified here (Figure S3). LARP4A and B version have similar conserved regions but with some significant differences in the precise organization of their La module\(^{[1]}\), in the amino acid sequence of their CR1 and CR2 regions (Figure S3) and in the size and primary amino acids conservation of their N-terminal regions (NTRs). LARP4A has a short unconserved region upstream of the PAM2 w, while LARP4B corresponding region is longer (designated by P2 + in Figure 3) and very well conserved (Figure S3). In a recent work, the human LARP4A NTR (position 1–111) was showed to adopt a semi-disordered state and function as RNA-binding platform, in combination with the RRM1 region\(^{[24]}\). The secondary structure propensity of this NTR and the conservation of the PAM2 w motif are both important determinants for RNA-binding. It would be interesting to test if differences observed in LARP4A and LARP4B NTRs reflect different RNA-binding properties of these two proteins.

**LARP6**

LARP6 genes are present basal Stramenopiles, in Chlorophytes, Embryophytes and Metazoans but are apparently absent from other eukaryotic lineages. Most Embryophytes possess three distinct LARP6 versions (green shading in Figure 4). For Metazoans, although invertebrates possess a single LARP6 version, most vertebrates have two (LARP6A and B, grey shading in Figure 4), with two exceptions: some fishes that have three versions (LARP6A, B1 and B2) and Eutherians that have only one (of the A type) (see Figure 4). All eukaryote LARP6 s present three highly conserved regions, the La motif and adjacent RRM (forming the LaM) and the C-terminal LAM and S1 Associated motif (LSA), that represent a specific diagnostic region of the LARP6 group. All Embryophyte LARP6s present an additional conserved GK/R-rich motif between the RRM and the LSA. Bony fish LARP6As also present a much longer N-terminal region compared to other LARP6s, but without any clear supplementary conserved regions.

**LARP7**

Full-length LARP7 s have a similar organization as genuine La proteins with a La motif, an RRM1 and an atypical RRM2. However, the spacer region between the LaM and the RRM2 is
much longer for LARP7 compared to genuine La, and the RRM2 is closer to the C-terminal end of the protein. LARP7s are present in all animals and in Amoebozoa (HaLARP7, DdLARP7), Alveolates (EaLARP7, SleLARP7, ThLARP7) and Stramenopiles (PsLARP7, TcLARP7) (Figure 5), but not all LARP7s belonging to these three later lineages host a conserved RRM2 region. Recently, several reports described the presence of a LARP7 (named Pof8) in fission yeast[10–12]. The only significant primary sequence conservation between Pof8 and other eukaryote LARP7s resides in the short RRM2 region, and the presence of a LaM (La motif/RRM1) can only be inferred from secondary sequence predictions. Accordingly, apart from the RRM2 region, primary sequence homology to Pof8 was not found outside of a few S. pombe sister species. Using a phylogenetic approach, we were able to confirm that the Pof8 RRM2 is indeed of the LARP7 type (and not of the genuine La type) (not shown). We next used Pof8 RRM2 as a bait, to detect proteins containing a similar RRM2 in species outside of the Taphrinomycotina lineage. Pof8-related RRM2 regions were detected in proteins from most Basidiomycota species. Surprisingly, except for species belonging to Pezizomycetes, no such protein could be detected in Ascomycota nor in other more basal Fungi lineages. These Pof8-related RRM2 regions are not only similar at the primary sequence level but also likely at the secondary structure level (see Figure 6). Some of these Pof8-like proteins are also predicted to adopt a LaM structure upstream of the RRM2 (not shown), suggesting that they could be a true orthologue of the S. pombe Pof8 protein. Others are however too short to host a LaM or are not predicted to fold as a LaM.

**Discussion**

This work provides new information on the detailed distribution and evolutionary history of La and LARPs in all major eukaryotic lineages. It also identifies several new conserved domains in genuine La and LARP1, 4 and 6 (see Figures 1 to 4 and S1 to S4) that could be the target of future functional studies.

One of the most surprising results of this study is the discovery that classical genuine La proteins are apparently absent from Alveolates, a major eukaryotic lineage. LARP7 proteins belonging to this lineage have been well characterized in two species (Tetrahymena thermophila and Euplotes aediculatus) [8,9], however, no genuine La from Alveolates have been reported so far. We find here that Alveolates possess an atypical La protein, with a conserved La motif, clearly grouping with classical genuine La motifs (see Figure 1). However, this ‘genuine La-like motif’ is not associated with RRMs but with a conserved motif of unknown function called DUF3223 (also named DeCI). DUF3223 is also detected in the C-terminal region of two plant-specific RNA Polymerases.
(Pol IV/V)[25], in three plant proteins likely involved in nuclear, chloroplastic and mitochondrial rRNA processing and ribosome assembly [26–28] and in many bacteria (Figure S1). The function of the DUF3223 is unknown, but this conserved region has been proposed to be involved in binding proteins and RNA, at least in the context of the plant PolV enzyme[25]. It is therefore tempting to speculate that these unusual La/DUF3223 containing proteins may represent the atypical Alveolates genuine La and that this combination of conserved regions may form a new and original RNA-binding surface. Functional studies will be needed to determine if La/DUF3223 proteins are able to bind RNA and fulfill classical genuine La functions in Alveolates.

Another intriguing aspect is the presence of two genuine La versions in two Embryophyte lineages (Brassicaceae and Poaceae) (see Figure 1). This situation likely results from two independent gene duplication events, one in the common ancestor of Poaceae (around 76MY ago), the other in the common ancestor of Brassicaceae (around 37MY ago). Although this second event is much younger than the previous one, Brassicaceae paralogues are much more divergent compared to the ones from Poaceae (see Figure 1). This suggests that a neofunctionalization event occurred in the Brassicaceae lineage that included a fast-evolving period. This hypothesis is supported by data showing that only one of the two Arabidopsis thaliana genuine La proteins is involved in tRNA maturation and is essential for viability[29]. It remains to be seen what the function of the second Brassicaceae La copy could be, and if the closely related Poaceae La paralogues are fully redundant or not.

The study of LARP1 confirmed the presence in all vertebrates of two distinct LARP1 versions (LARP1A and B), a situation previously described in human. All works describing LARP1 function in mammals have been done on LARP1A and the function of the LARP1B version in vertebrates remains to be studied. An intriguing aspect of LARP1 evolution is the presence in two lineages (Fungi and Embryophytes) of proteins containing a solo La motif (of the LARP1 type) in a C-terminal position. Saccharomyces cerevisiae bears two of these solo La motif LARP1, named Sfl1 and Sro9p, that are both involved in the regulation of mRNA localization, stability and translation.
The La motif of Slf1 and Sro9p is necessary but not sufficient to bind RNAs, and the additional cis and/or trans factor(s) needed to generate the required RNA-binding surface have not been detected so far[30]. In Embryophyte genomes, solo La motif LARP1 are systematically present in association with classical full-length LARP1. LARP1 distribution in Embryophytes and Fungi suggests that solo La motif LARP1 versions appeared two times independently during eukaryote evolution.

Figure 5. Distribution, phylogenetic relationships and structural organization of LARP7 in eukaryotes. Full-length LARP7 proteins from the different eukaryotic lineages were aligned and used to construct the phylogenetic tree. The Fungi LARP7-related (Pof8) proteins were not used for the phylogenetic analysis since their homology with other LARP7 is limited to the short RRM2 region. Fungi LARP7-like La motif and RRM1 were only inferred from secondary structure predictions and are represented by La* and RRM1* in the corresponding cartoon. Scale bar indicates length of 1 substitution/site. For a description of species and protein sequences used, see Supplemental Table 1. Boxes on the right define how the different lineages were colour-coded. Positions of the different conserved regions (on the Human LARP7 sequence (NP_056269)) are the following: La (40–111), RRM1 (124–199), RRM2 (450–558).

Figure 6. Sequence alignment of Pof8 (LARP7-like) RRM2 from different Fungi lineages showing primary and secondary sequence conservation. The amino acids are coloured in blue based on their conservation. Experimentally determined positions of the S. Pombe Pof8 (SpPof8) secondary structure[39] are schematically represented above the alignment. For other sequences, secondary structure predictions, inferred using the Phyre2 software are superimposed. The origin of each sequence is colour-coded.
evolution, once in the common ancestor of Angiosperms and once in the common ancestor of Dikarya (at the origin of Basidiomycota and Ascomycota). While in Angiosperms, both full-length and solo La motif versions were kept in all species, in Dikarya the solo version is the rule, as full-length LARP1 were most often lost or largely reorganized (in Pezizomycotina). It remains to be seen if Embryophytes and Fungi solo La motif LARP1s have similar functions (i.e. convergent evolution). Also, the rapid divergence of full-length LARP1 in Pezizomycotina could be symptomatic of a neofunctionalization event. It would therefore be interesting to determine if these more divergent proteins are still fulfilling the same functions as other full-length eukaryotic LARP1s.

The evolutionary history of LARP4 and LARP6 is more complex compared to the ones of genuine La and other LARPs (the two evolutionary scenarios described below are summarized in Figure 7). For LARP4, its unusual distribution in eukaryotes suggests the following evolutionary scenario (Figure 7A). A LARP4 gene was present very early in eukaryote evolution, as suggested by its presence today in some basal eukaryotic lineages (Excavates, Amoebozoa, Stramenopiles). However, contrarily to LARP1, LARP4 gene was lost in Embryophytes, Chlorophytes and Fungi. A first gene duplication event occurred in the common ancestor of vertebrates generating the two LARP4 versions (A and B). A second gene duplication, targeting the LARP4A version, occurred in fishes but not at the onset of this class (around 500MY ago) but later in the common ancestor of the Clupeocephala group (around 250MY ago). This results in cartilaginous fishes (Chondrichthyes) and some bony fishes belonging to basal orders (such as Coelacanthiformes, Polypteriformes, Acipenseriformes, Lepisosteiformes and Osteoglossomorpha) having a single LARP4A gene while other fishes have two (see Figures 3 and 7).

The evolutionary history of the LARP6 family is also complex and includes several lineage-independent gains and losses (see Figures 4 and 7B). A LARP6 gene was also likely present early in eukaryote evolution, as suggested by its presence today in Stramenopiles. In Embryophytes, in the common ancestor of Angiosperms, gene duplication events led to the emergence of three LARP6 versions (A, B and C). These three versions are systematically present in all modern Angiosperm species. The LARP6A version is always encoded by a single copy gene, but LARP6B and C versions are sometimes encoded by multiple paralogous genes. In Metazoans, while invertebrates possess a single LARP6 gene, a gene duplication event (independent from the one proposed in Embryophytes) likely occurred in the common ancestor of vertebrates, generating LARP6A and B types. Vertebrates LARP6B type was further duplicated in fishes, in the common ancestor of Actinopterygii (around 375 MY), to generate the B1 and B2 types. However, the LARP6B2 type was lost two times in evolution, once precisely in Cypriniformes (that include the model species Danio rerio), and a second time in the common ancestor of Neoteleostei (around 180 MY). Therefore, Cypriniformes and Neoteleostei present only a single LARP6B of the B1 type. The loss of the LARP6B2 type in the common ancestor of Neoteleostei was apparently associated with the divergence of the remaining LARP6B (see the longer branch length for AteLARP6B, LbLARP6B, LcaLARP6B, BpLARP6B and OiLARP6B on the tree of Figure 4 compared to other fish LARP6 s). Another independent gene loss, this time of the LARP6B type, occurred in the common ancestor of Eutherians, leaving a single LARP6 (of the A type) in these species.

Recently, a LARP7-like protein (Pof8) was described in Schizosaccharomyces pombe and in a few Taphrinomycotina sister species [10–12]. Primary sequence homology of Pof8 to other eukaryotic LARP7s is limited to the RRM2 region and consequently, Pof8 LaM was only inferred from secondary structure predictions. Although most LARP1 proteins possess RRM1 regions conserved only at the secondary sequence level, this is
the first time that a La-related protein presents only secondary structure conservation of the complete LaM. This suggests that some functions fulfilled by LARP7s require only partial (most LARP1s) or no (Pof8) primary conservation of the LaM, while still needing the correct folding of this region. Using the S. pombe pof8 RRM2 sequence, we were able to identify several Pof8-related proteins in three other Fungi lineages (see Supplemental Table 1 and Figure 6). Pof8-related proteins all possess primary and likely secondary structure homologies to the S. pombe RRM2 (see Figure 6). However, some of these Pof8-related proteins are too short to host a LaM or were not predicted to fold as a LaM. In these cases, one possibility would be that they could interact in trans with a yet to identify partner to provide the missing structures. The patchy distribution of Pof8-like proteins in Fungi still remains difficult to explain. Indeed, although Pof8-like proteins are very well represented in Basidiomycota, they are absent from most Ascomycota lineages (except for Taphrinomycotina and Pezizomycetes) and from basal Fungi lineages. Since Pof8 is involved in telomerase assembly[10–12], the loss of Pof8-like proteins may indicate functional differences in the way telomerase associates with its telomerase RNA in these Pof8-free lineages.

Methods

Blast searches (blastp and tblastn) were performed starting from known La and LARP7s conserved domains on the genome of 135 species representing all major eukaryotic lineages using the JGI (https://jgi.doe.gov) and NCBI (https://www.ncbi.nlm.nih.gov) genomic resources. Full-length protein sequences were aligned using the multiple sequence comparison by log-expectation (MUSCLE v3.7) software[33]. The resulting alignment was processed using the TRIMAl software [34] and trees were reconstructed using the fast maximum likelihood tree estimation program PHYML [35] using the LG amino acids replacement matrix [36]. Statistical support for the major clusters was obtained using the approximate likelihood-ratio test (aLRT) [37]. Protein secondary structure predictions were obtained using the Phyre2 (V2) software[38].

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Disclosure statement

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