

# Convergent origins and rapid evolution of spliced leader *trans*-splicing in Metazoa: Insights from the Ctenophora and Hydrozoa

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## ABSTRACT

Replacement of mRNA 5' UTR sequences by short sequences *trans*-spliced from specialized, noncoding, spliced leader (SL) RNAs is an enigmatic phenomenon, occurring in a set of distantly related animal groups including urochordates, nematodes, flatworms, and hydra, as well as in Euglenozoa and dinoflagellates. Whether SL *trans*-splicing has a common evolutionary origin and biological function among different organisms remains unclear. We have undertaken a systematic identification of SL exons in cDNA sequence data sets from non-bilaterian metazoan species and their closest unicellular relatives. SL exons were identified in ctenophores and in hydrozoan cnidarians, but not in other cnidarians, placozoans, or sponges, or in animal unicellular relatives. Mapping of SL absence/presence obtained from this and previous studies onto current phylogenetic trees favors an evolutionary scenario involving multiple origins for SLs during eumetazoan evolution rather than loss from a common ancestor. In both ctenophore and hydrozoan species, multiple SL sequences were identified, showing high sequence diversity. Detailed analysis of a large data set generated for the hydrozoan *Clytia hemisphaerica* revealed *trans*-splicing of given mRNAs by multiple alternative SLs. No evidence was found for a common identity of *trans*-spliced mRNAs between different hydrozoans. One feature found specifically to characterize SL-spliced mRNAs in hydrozoans, however, was a marked adenosine enrichment immediately 3' of the SL acceptor splice site. Our findings of high sequence divergence and apparently indiscriminate use of SLs in hydrozoans, along with recent findings in other taxa, indicate that SL genes have evolved rapidly in parallel in diverse animal groups, with constraint on SL exon sequence evolution being apparently rare.

**Keywords:** *trans*-splicing; spliced leader; evolution; *Clytia*; *Pleurobrachia*

## INTRODUCTION

The process of spliced leader (SL) *trans*-splicing joins the short exon (15–50 nucleotides [nt]) from specialized noncoding nuclear SL RNAs to the 5' ends of assorted pre-mRNAs during their nuclear processing (for reviews, see Davis 1996; Hastings 2005). This unusual form of RNA maturation was first described in trypanosomes, in which all mRNAs are *trans*-spliced, using a single SL RNA (Sutton

and Boothroyd 1986). SL *trans*-splicing has subsequently been demonstrated to occur in Euglenozoa (Miller et al. 1986; Tessier et al. 1991), in dinoflagellates (Zhang et al. 2007; Zhang and Lin 2009), and in several metazoan lineages: all tested nematodes (Krause and Hirsh 1987; Guiliano and Blaxter 2006), urochordates (Vandenbergh et al. 2001; Ganot et al. 2004), flatworms (Davis 1997), the two tested chaetognath species (Marletaz et al. 2008), a Bdelloid rotifer (Pouchkina-Stantcheva and Tunnacliffe 2005), an Acoel (Marletaz et al. 2008), and the hydrozoan (Cnidaria) *Hydra vulgaris* (Stover and Steele 2001). In contrast, SL *trans*-splicing was not detected in vertebrates, echinoderms, arthropods, mollusks, or annelids, or in most nonmetazoan groups including fungi and plants. Furthermore, there are wide differences in SL usage within and between different animal groups, such as in the proportion

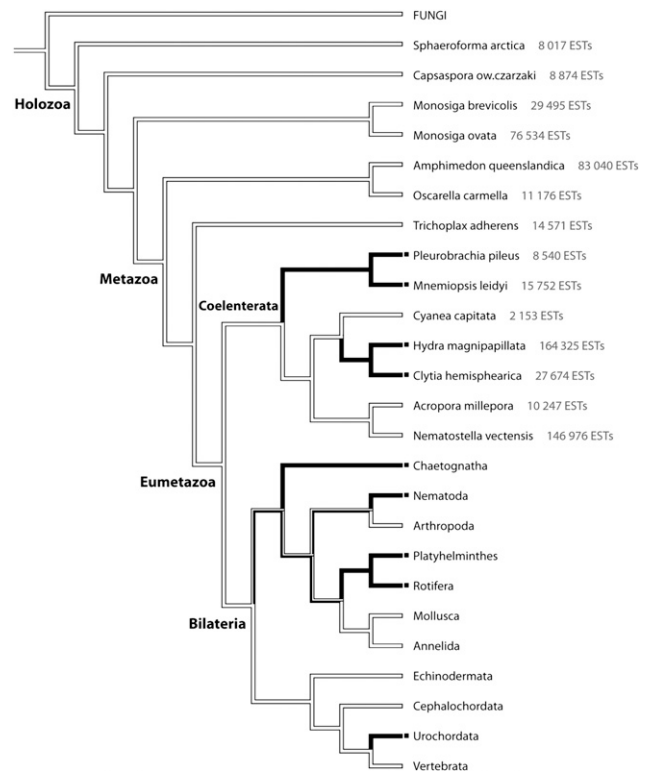
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of *trans*-spliced mRNA and the number of distinct SL genes (for reviews, see Hastings 2005; Marletaz et al. 2008).

SL *trans*-splicing is enigmatic not only because of its scattered phylogenetic distribution and variable frequency, but because of the lack of a clear conserved function. In all cases in trypanosomes, and in some nematodes, urochordates, chaetognaths, and possibly flatworms, SL *trans*-splicing has an important function in the resolution of polycistronic pre-mRNAs into individual transcripts (Davis and Hodgson 1997; Blumenthal and Gleason 2003; Satou et al. 2006; Marletaz et al. 2008). In kinetoplastids (i.e., trypanosomes and Euglenozoas) and nematodes (Lall et al. 2004), SL *trans*-splicing affects stability and translation by providing a particular 5' cap structure, the kinetoplastid case being extreme in that the new cap is required for association with ribosomes (Zeiner et al. 2003). It has also been pointed out that the shortening of 5' UTRs that results from SL *trans*-splicing could potentially act to "sanitize" overlong pre-mRNAs by removing unwanted sequences potentially affecting translation or stability from the original "outtron" (i.e., the part of the original 5' UTR replaced by the SL sequence), thereby facilitating the use of more distant transcription initiation sites (Davis 1996; Hastings 2005).

Concerning the evolution of SL genes, available evidence has not allowed discrimination between hypotheses of a common origin with multiple losses and independent acquisition in multiple phylogenetic lineages (Lawrence 1999; Nilsen 2001; Stover and Steele 2001; Hastings 2005; Roy and Irimia 2009). Arguments put forward to favor the former hypothesis include the existence of (partially) shared functions, such as polycistronic transcript resolution, between some distantly related taxa, the stereotypic secondary structure of the SL RNA "intron" region, which features three stem-loops and a binding site for the Sm proteins (essential components of the eukaryotic *cis*-splicing machinery), and the frequent localization of SL genes within 5S rRNA gene clusters. However, given the phylogenetic distribution of species known to employ SL *trans*-splicing among metazoans, convergent acquisitions of structural and functional features following multiple independent origins cannot be excluded. In order to shed light on SL evolution in metazoans, and, in particular, on their presence or absence in the common metazoan ancestor, we have undertaken a systematic search for SL exons among expressed sequence tag (EST) data sets of early diverging metazoans (cnidarians, ctenophores, placozoa, and sponges) and in four species of unicellular organisms that group together with the metazoans in the clade Holozoa (Fig. 1). We then performed a detailed comparison of SL use in two hydrozoan species, *Clytia hemisphaerica* and *Hydra magnipapillata*, since short-range comparisons can be very informative about conserved features and sequence divergence rate, as shown by studies in the nematodes and chaetognaths (Guiliano and Blaxter 2006; Marletaz et al. 2008).



**FIGURE 1.** SL usage in holozoan species. The character "presence/absence" of SL genes, deduced from this study (with numbers of ESTs analyzed per species) and from published work, mapped using Mesquite parsimony reconstruction (Maddison and Maddison 2009) onto the phylogeny of Holozoa. This phylogeny was adapted from Dunn et al. (2008) for intrabilaterian relationships and from Philippe et al. (2009) for all other relationships. All SL-positive lineages found in the Metazoa have been included with the exception of the Acoels, whose phylogenetic position remains unclear (Dunn et al. 2008; Hejnol et al. 2009; Schierwater et al. 2009). (Black lines) Presence of SL genes, (white lines) their absence, (mixed lines) unresolved state. The rotifers have been coded as "unresolved" because current evidence indicates SL usage in bdelloid but not monogonont rotifer species (Pouchkina-Stancheva and Tunnacliffe 2005; Suga et al. 2007). While this study was under review, SL sequences were also discovered in a Hexactinellid sponge and a third ctenophore species (Douris et al. 2009), not included here.

## RESULTS

### Survey of SL *trans*-splicing in holozoans from EST data sets

The presence of an identical sequence at the 5' end of *trans*-spliced mRNAs means that SL *trans*-splicing can be readily identified in large EST data sets by looking for identical sequences in the 5' UTRs of assembled cDNAs. Large EST collections were generated for the ctenophore *Pleurobrachia pileus* and the hydrozoan cnidarian *Clytia hemisphaerica* at the Genoscope (Evry, France) using mixed-stage cDNA libraries (Philippe et al. 2009). EST sets for 12 other species were downloaded from NCBI: four additional cnidarians (the Hydrozoan *Hydra magnipapillata*, the Anthozoans

*Nematostella vectensis* and *Acropora millepora*, and the Scyphozoan *Cyanea capillata*), the ctenophore *Mnemiopsis leidyi*, the sponges *Oscarella carmela* (Homoscleromorpha) and *Amphimedon queenslandica* (Demospongia), the placozoan *Trichoplax adherens*, the choanoflagellates *Monosiga brevicollis* and *Monosiga ovata*, the ichthyosporean *Sphaerofoma arctica*, and the related ameboid protist *Capsaspora owczarzaki*. For each data set, cDNA sequences were assembled from the ESTs, and identical or nearly identical 5' leader sequences common to multiple transcripts were identified following alignment (see Materials and Methods for details).

Using this exon detection approach we could clearly identify 5' SL exon sequences in EST assemblies from the two ctenophore species, *P. pileus* and *M. leidyi*, and in the two hydrozoans, *C. hemisphaerica* and *H. magnipapillata* (Fig. 1). We were also able to identify several SL sequences from the small EST data set of a third hydrozoan species, *Podocoryne carnea* (3905 ESTs, downloaded from The National Center for Biotechnology Information [NCBI]; <http://www.ncbi.nlm.nih.gov/guide/>) (data not shown). In contrast, no conserved 5' elements were detected in the EST data set from any of the other species tested, strongly suggesting that SL *trans*-splicing is absent in these species (although we cannot completely rule out the possibility of unprecedentedly low SL *trans*-splicing frequencies). In each data set from which putative SL exons were identified, many sequence variants were found, defined by their presence in at least five distinct cDNA assemblies (see below for detailed analyses). We confirmed that each variant identified in *H. magnipapillata* corresponded to

a distinct SL gene by BLAST analysis against the draft genome of this species (<http://hydrazome.metazome.net>); all variants were present in the genome sequence with the exception of one SLB variant.

To see whether our new data could help clarify the picture of SL evolution, we mapped the presence or absence of SL genes obtained from our analyses onto a phylogenetic tree combining recent topologies obtained by phylogenomics approaches focusing on basal metazoan branching (Philippe et al. 2009) and on intrabilaterian relationships (Dunn et al. 2008). SL sequences appear to be restricted to a small number of lineages among Eumetazoa: Ctenophora, Hydrozoa, Urochordata, and several protostome lineages. Parsimony optimization of SL evolution fails to resolve the ancestral state of Protostomia but clearly supports absence of SL *trans*-splicing in both metazoan and eumetazoan ancestors, followed by multiple independent origins (Fig. 1).

### Multiple spliced leaders in ctenophores

Our results provide the first evidence for SL *trans*-splicing in ctenophores, suggesting that SL usage within the Coelenterate lineage has been acquired (or lost) at least twice (Fig. 1). As in the hydrozoans (Stover and Steele 2001 and see below), multiple SL sequences were recovered from *P. pileus*. SL exons were detected in 40% of the ~9000 distinct assembled cDNA sequences. These sequences could be grouped into two distinct but related groups, Ppi\_SLA and Ppi\_SLB, with a maximum length of 37 nt recovered in both groups (Table 1). Ppi\_SLA showed nine distinct variants

**TABLE 1.** SL sequences from the ctenophores *Pleurobrachia pileus* and *Mnemiopsis leidyi*

Species	SL name	SL sequence	<i>Trans</i> -spliced cDNAs (%)
<i>Pleurobrachia pileus</i>			
	Ppi_SLA1	CUUUUC-ACACUACUUUAAACAAAUUAAUU-UGAG	77
	Ppi_SLA2	AACUUUUAACACUACUUUAAACAAAUUAAUU-UGAG	
	Ppi_SLA3	CUUUUC-ACACUACUUUAAACAAAUUAAUU-UGAG	
	Ppi_SLA4	CACUACUUUAAACAAACUAAUU-UGAG	
	Ppi_SLA5	CUACUUUAAACAAAUUAAUUUUGAG	
	Ppi_SLA6	CACUACUUUAAACAAAUUAAUUUUGAG	
	Ppi_SLA7	CUUUAAACAAAUUAAUU-UGAG	
	Ppi_SLA8	AACUUUUAACACUACUUUAAACAAAUUAAUUUUGAG	
	Ppi_SLA9	CUACUUUAAACCA-UUAAUU-UGAG	
	Ppi_SLB1	AACUUUCAUACAACUACAACGUAACAAUUAUUUUGAG	23
	Ppi_SLB2	AACUUUCAUACAACUACCACGUAACAAUUAUUUUGAG	
<i>Mnemiopsis leidyi</i>			
	Mle_SLA	ACUUUCAACACUACUAAUUAACAAAUUAAUUUGAG	47.7
	Mle_SLB	UUUUAAUACUUUCAACAACUACUAAUUAUUAAUUUUGAG	16.3
	Mle_SLC	UUUUAAACUAAUUUCAACUACAACAAAUUAAUUUUGAG	18.9
	Mle_SLD	CUACAAAUAUCAAAUUUUAUUUGAG	17.4

List of SL variants detected in *P. pileus* and *M. leidyi* ESTs. SL sequences are aligned within each SL group.

(probably representing nine distinct genes, see above) and was detected in 77% of *trans*-spliced cDNAs, while Ppi\_SLB had two variants. The Ppi\_SL exon sequences obtained were mostly incomplete in their 5' ends, but the longest variants of each groups exhibited a common 5' terminal motif, AAC(U)<sub>n</sub>CA. We found clear evidence that given pre-mRNAs can be joined to alternative SLs, with 103 distinct transcripts in the data set found to be *trans*-spliced either by Ppi\_SLA or Ppi\_SLB groups. Among these multiple *trans*-spliced cDNAs, 12 showed different acceptor splice sites for Ppi\_SLA and Ppi\_SLB, whereas in all others splicing was at exactly the same position in the 5' UTR (see the examples in Supplemental Data S1).

Multiple SL sequences were also recovered from EST data of another ctenophore species, *M. leidy*. Four different SL groups (Table 1, Mle\_SLA, Mle\_SLB, Mle\_SLC, Mle\_SLD), each with only one variant, were found in 3% of the 2968 assembled cDNAs. In terms of sequence similarity, all four Mle\_SL groups are closely related to each other but different from the Ppi\_SL variants. The absence of identical sequences between these two ctenophore species was confirmed by a negative search of Mle\_SL sequences in *P. pileus* ESTs and of Ppi\_SL sequences in *M. leidy* ESTs.

### High diversity of spliced leader groups in hydrozoans

Analysis of data sets for the hydrozoans *H. magnipapillata* and *C. hemisphaerica* revealed a high number of SL sequence variants per species. Given the *H. magnipapillata* genome analysis (see above), we assume that all of these variants correspond to distinct SL genes. It should be emphasized that the diversity in SL sequences is probably underestimated due to selective transcriptome representation and incomplete 5' termini in the assembled ESTs. In the most complete EST data set, that of *C. hemisphaerica*, SL exons were detected in 23% of the approximately 19,000 distinct assembled cDNA sequences. SL sequence variety in *C. hemisphaerica* was even greater than that detected in ctenophore species, with five distinct groups of SL exon sequences (Table 2, Che\_SLA to Che\_SLE). Each SL sequence group showed several variants (putative genes), with the exception of Che\_SLA, which despite being represented in >20% of *trans*-spliced cDNA assemblies showed only a single variant. Che\_SLB group exons were detected most frequently, in over half of *trans*-spliced cDNA assemblies, while Che\_SLC and Che\_SLE exons were rare. The *trans*-splicing frequency for particular SL groups thus did not correlate with the number of member SL genes. We could not detect a common 5' motif of Che\_SL sequences, perhaps because most 5' termini are incomplete in the assembled ESTs.

The *H. magnipapillata* EST set showed a similar overall pattern of SL use, with six spliced leader groups and a total of 15 variants detected among ~3000 of 25,000 assembled cDNA sequences that showed *trans*-splicing. The Hma\_SL groups were again found to be distinct but related, with

a common 5' motif ACGG(A)<sub>n</sub>C detectable in all six groups (the 5' end of *Hydra* SL sequences were completed using genomic data) (Table 2). The relatively low percentage (12%) of *trans*-spliced sequences detected in the *Hydra* versus *Clytia* transcriptome data sets likely reflects in part differences in the origins or qualities of the cDNA libraries used for EST sequencing.

One SL group was detected in nearly 80% of *trans*-spliced *H. magnipapillata* cDNAs, and was designated Hma\_SLB because of its 100% identical nucleotide sequence with the previously characterized SLB from *H. vulgaris* (Stover and Steele 2001). No sequence identical to the *H. vulgaris* SLA exon was detected in the *H. magnipapillata* cDNA data set and genome; however, studies of genomic DNA revealed that *H. vulgaris* SLA corresponds to a sequence we designated Hma\_SLA1, despite the low similarity of the two sequences (see below). Reverse searching of *H. vulgaris* EST data revealed the presence of most Hma\_SLB and Hma\_SLC variants, previously unreported, with SLB group exons again detected in the majority, indicating that most of the multiple SL genes are shared between these closely related *Hydra* species.

The absence of identical SL sequence between *Hydra* and *Clytia* was confirmed by negative BLAST searches for *C. hemisphaerica* SL sequences in the *H. magnipapillata* draft genome and for *Hydra* SL sequences in the *Clytia* ESTs. Although the sequences of SLs from different hydrozoan species (as well as between hydrozoan and ctenophoran SL exons) may well be evolutionarily related, the lack of sequence similarity between them was so great that it precluded phylogenetic analysis to evaluate their evolutionary relationships.

### Rapid SL evolution at the genomic level in hydrozoans

The evidence for rapid SL gene evolution obtained from analysis of SL representation in the transcriptome was extended by comparison of two SL gene sequences and the surrounding genomic regions between *Hydra* species. A previous study in *H. vulgaris* revealed a spliced leader gene in each of two inter-5S rRNA gene regions amplified by PCR (Stover and Steele 2001). We aligned these with equivalent regions identified by BLAST from *H. magnipapillata* genome sequences. One of the regions contains the Hma\_SLB1 gene in *H. magnipapillata* and its direct counterpart in *H. vulgaris* (Fig. 2A). The SLB1 exon is perfectly conserved between the two species, while the intron domain shows one difference per 10 nucleotides (Fig. 2B).

In contrast, the other inter-5S region provided evidence of clear SL gene diversification between the two *Hydra* species, containing the sequences defined as Hma\_SLA1 in *H. magnipapillata* and as SLA in *H. vulgaris*. The identical genomic position and orientation of these clearly distinct SL sequences within this 5S rRNA gene cluster suggests a common origin from an ancestral SLA gene. The exon of this SLA gene has clearly evolved faster than the intron domain

**TABLE 2.** SL sequences from the hydrozoans *Clytia hemisphaerica* and *Hydra magnipapillata*

Species	SL name	SL sequence	Trans-spliced cDNAs (%)
<i>Clytia hemisphaerica</i>			
	Che_SLA	AAAAAAUUCACUCCAUAAGAUAUAGUGAAUAAG	23.6
	Che_SLB1	GAAAAAAUACUC-AUUCCAUUUAGUCACUGAGUAUAAG	56.1
	Che_SLB2	AAAAUACUC-AUUUCAUUUAGUCACUGAGUAUAAG	
	Che_SLB3	AAAUACUCUAAUCCAUUUAGUCACUGAGUAUAAG	
	Che_SLB4	AAAAUACUCUAAUUAAUUUAGUCACUGAGUAUAAG	
	Che_SLB5	CUCUAAUUAAUUUAGUUACUGAGUAUAAG	
	Che_SLB6	AAAAUACUC-AUUCCAUU-AGUCACUGAGUAUAAG	
	Che_SLB7	AAAAUACUC-AUUCCAUUCAGUCACUGAGUAUAAG	
	Che_SLB8	UACUC-ACUCCAUUUAGUCACUGAGUAUAAG	
	Che_SLB9	AAAAUACUC-AUUCCAUUUAGUUACUGAGUAUAAG	
	Che_SLB10	AAAAUACUC-AUUCCAUUUAGUCACUGAGUCUAAG	
	Che_SLB11	AUUCCAUUUAGUCACUGAGUUUAAG	
	Che_SLC1	GAAACACCUUUCUAGACUCUAUUUAGUUGAGCAUAAG	1.2
	Che_SLC2	GAAACUCCUUUCUAGAUUCUAUUUAGUUGAGCAUAAG	
	Che_SLC3	GAAACACCAUUCUAGACUCUAUUUAGUUGAGCAUAAG	
	Che_SLC4	GAAACACCUUUCUAGACUCUAUUUAGUUGAGCAUAAG	
	Che_SLD1	GAAACACCAUUACUCUAAAUUUGAGUCUAAG	17.4
	Che_SLD2	AGAAACACC-AUAAUCUCCAAUUUUGAGUCAAAAG	
	Che_SLD3	AGAAACACCAUUACUCUCCAAUUUUGAGUUUAAG	
	Che_SLD4	AACACCAUUACUCUAAAUUUGAGUUUAAG	
	Che_SLD5	AGAAACACCAUUAAUUCAAUUUUGAGAUUAAG	
	Che_SLD6	AGAAACACCAUUACUCUCCAAUUUUGAGUAUAAG	
	Che_SLD7	AGAAACACCAUUACUCUAAAUUUGAGUAUAAG	
	Che_SLE1	AAAAACACCAUACUCUCCUGAUUUUUGACUACAAG	1.7
	Che_SLE2	AAAAACCAUACUCUCCUGAUUUUUGACUAAAAG	
	Che_SLE3	AAUCACUCCUGAUUUUUGAGUAUAAG	
<i>Hydra magnipapillata</i>			
	Hma_SLA1	ACGGAAAAAACACAAUAAACAAACAGUUCUAUUUUGUGUUAUAAG	11.5
	Hma_SLA2	ACGGAAAAAACACAAUAAACAAACAGUUCUAUUUUGUGUUAUAAG	
	Hma_SLB1 <sup>a</sup>	ACGGAAAAAACACAUACUGAAACUUUUUAGUCCUGUGUAAUAAG	78.2
	Hma_SLB2 <sup>a</sup>	ACGGAAAAAACACAUACUGAAACUUUUUAGUCUUUGUGUAAUAAG	
	Hma_SLB3 <sup>a</sup>	ACGGAAAAAACACAUACUGAAACUUUUUAGUCCUGUGUAAUAAG	
	Hma_SLB4 <sup>a</sup>	ACGGAAAAAACACAUACUGAAACUUUUUAGUCCUUUGUAAUAAG	
	Hma_SLB5 <sup>a</sup>	ACGGAAAAAACACAUACUGAAACUUUUUAGUCUCUGUGUAAUAAG	
	Hma_SLB6 <sup>a</sup>	ACGGAAAAAACACAUACUGAAACUUUUUAGUCCUGUGUAAUAAG	
	Hma_SLB7	ACGGAAAAAACACAUACUAAAACUUUUUAGUCCUGUGUAAUAAG	
	Hma_SLC <sup>a</sup>	ACGGAAAAACGCAUUUAUAAACUCUUGUUUUAUUGCGUAAUAAG	3
	Hma_SLD	ACGGAAAAAACACAAACAAAACUCAACGUUAAAUUUGUGUAAUAAG	2.4
	Hma_SLE1	ACGGAAAAAACACAUCAAAACUUUUUUUAGUAUUUGUGUCAUAUAAG	4.3
	Hma_SLE2	ACGGAAAAAACACAUCAAAACUUUUAAGUAUUUGUGUCAUAUAAG	
	Hma_SLE3	ACGGAAAAAACACAUCAAAACUUUUUUUAGUAUUUGUGUCAUAUAAG	

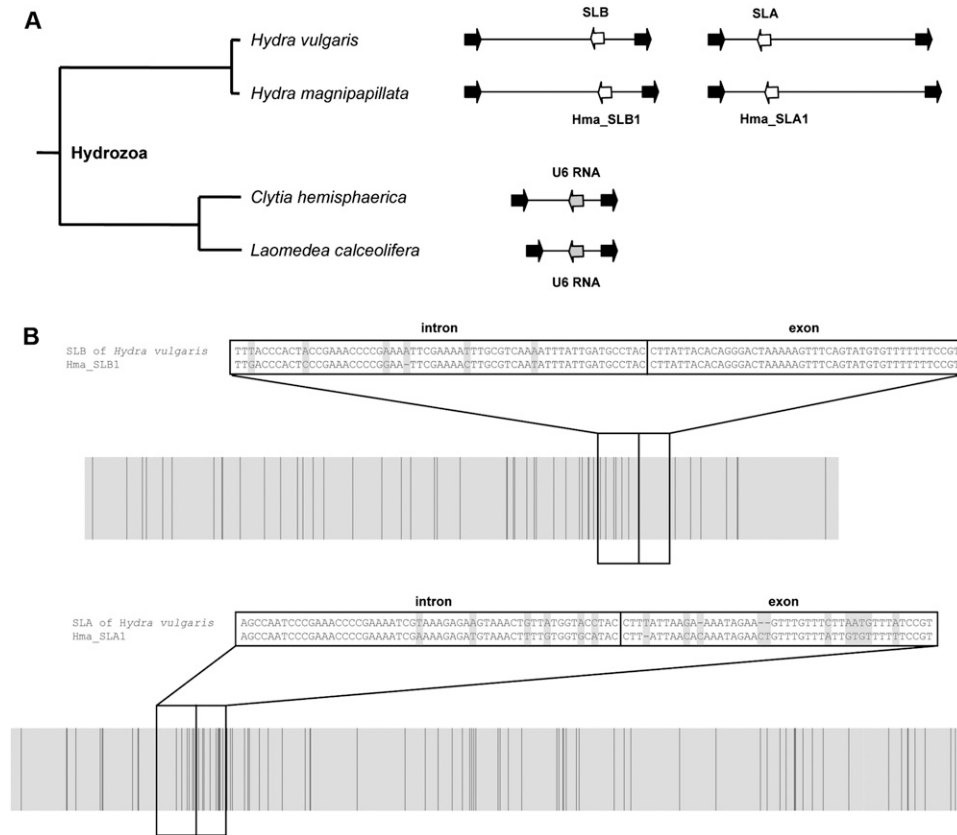
SL variants detected in *C. hemisphaerica* and *H. magnipapillata* ESTs. SL sequences are aligned within each SL group.

<sup>a</sup>Hma\_SL variants recovered as identical by BLAST in *H. vulgaris* ESTs.

(2.2 differences per 10 nucleotides and 1.1 differences per 10 nucleotides, for exon and intron domains, respectively); generally, the intron domain shows more differences than the exon domain (as observed for the SLB gene). It is also noteworthy that the SL genes evolved at least as fast as the surrounding 5S intergenic regions (1.6 differences per 10 nucleotides, 0.6 differences per 10 nucleotides, and 0.5 differ-

ences per 10 nucleotides, for the SLA gene, SLB gene, and 5S intergenic regions without SL genes, respectively). Although limited, these genomic sequence comparisons provide evidence for rapid SL gene evolution in *Hydra*, with the exon domains showing different degrees of divergence.

PCR amplification of inter-5S gene regions from genomic DNA of *C. hemisphaerica* and another hydrozoan



**FIGURE 2.** Identification of hydrozoan SL genes. (A) Small RNA genes found in between 5S rRNA genes (black arrows) in hydrozoans. (B) Alignment of *H. vulgaris* and *H. magnipapillata* regions. (Light gray) Conserved positions, (darker bars) mutations (each indel was treated as one mutation, independent of length). Sequences of orthologous SL genes (intron and exon) encoded by this genomic regions are shown.

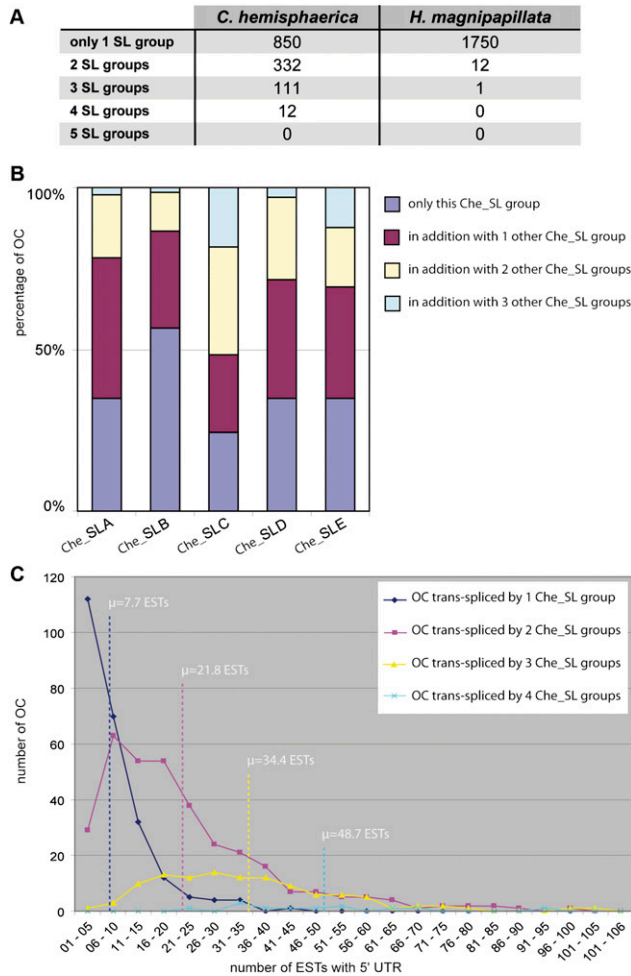
belonging to the same family (Campanulariidae), *Laomedea calceolifera*, yielded a single equivalent genomic fragment flanked by 5S genes (Fig. 2A), but no *Clytia* SL sequences were identified. In both species, the recovered inter-5S region housed a highly conserved U6 RNA gene in the same orientation (Fig. 2A; Supplemental Data S2). In contrast, in analysis of the *H. magnipapillata* genome, the multiple copies of U6 RNA genes identified by BLAST did not show an association with 5S rRNA genes, being positioned in different assembled scaffolds. It is clearly not possible to draw strong conclusions about SL evolution at the genomic level from the limited examples examined here; however, our findings are consistent with the marked plasticity in the linkage between SL and 5S ribosomal genes well demonstrated in the nematodes (Drouin and de Sa 1995).

### Indiscriminate SL *trans*-splicing in hydrozoans

The availability of a large transcriptome data set for *C. hemisphaerica* and *H. magnipapillata* allowed us to address whether particular SL exons showed any qualitative preferences in *trans*-splicing within and/or between species. We first assigned each *trans*-spliced cDNA sequence from *C. hemisphaerica* and *H. magnipapillata* to putative cnidarian

“orthology clusters” (OCs) (see Materials and Methods), each of them representing a single common gene. As previously noted for HTK32 and Syk genes in *H. vulgaris* (Stover and Steele 2001), many OCs in our data set showed *trans*-splicing by different SL groups, with 35% of *C. hemisphaerica* OCs being *trans*-spliced by more than one SL group (Fig. 3A; Supplemental Data S3). *Trans*-splicing of given mRNAs to alternative SLs was also detectable in the *H. magnipapillata* data set, albeit in only 1% of OCs (Fig. 3A).

The data set of 2643 *trans*-spliced OCs used in this analysis was assembled from 2967 cDNA sequences from *C. hemisphaerica* and 3054 from *H. magnipapillata*. Of these OCs, 880 contained *C. hemisphaerica* but not *H. magnipapillata* sequences, 1338 only *H. magnipapillata* sequences, and 425 sequences from both species (Fig. 4). The representation of each SL group in this 2643 OC data set corresponded to that in the original cDNA sequence collection. Among the 425 OCs *trans*-spliced in both species there was no preference for particular Che\_SL groups and Hma\_SL groups to associate with common transcripts, or for particular Che\_SLs to do so among the 455 OCs showing *trans*-splicing to more than one SL. In both cases, the SL groups were found proportionally represented among *trans*-spliced OCs (data not shown).



**FIGURE 3.** Indiscriminate SL *trans*-splicing in Hydrozoa. (A) Number of orthologous clusters (OCs) *trans*-spliced by one, two, three, four, and five SL groups identified in the *C. hemisphaerica* and *H. magnipapillata* OC data set. (B) Percentage of OCs in which a given Che\_SL group is found solely with or in addition to other Che\_SL groups among *C. hemisphaerica* cDNAs. The numbers of OCs analyzed here are 617, 884, 33, 412, and 49 for Che\_SLA, Che\_SLB, Che\_SLC, Che\_SLD, and Che\_SLE, respectively. (C) Number of ESTs with a 5' UTR in OCs *trans*-spliced in *C. hemisphaerica* cDNAs by one, two, three, and four SL groups. For each OC subset, the mean number of ESTs with a 5' UTR is given ( $\mu$ , dashed line).

Moreover, examples were recovered of individual OCs *trans*-spliced by one of any of the five Che\_SL groups and one, two, or three of the other groups (Fig. 3B). Taken together, these observations strongly argue in favor of indiscriminate use of all SL groups in hydrozoans.

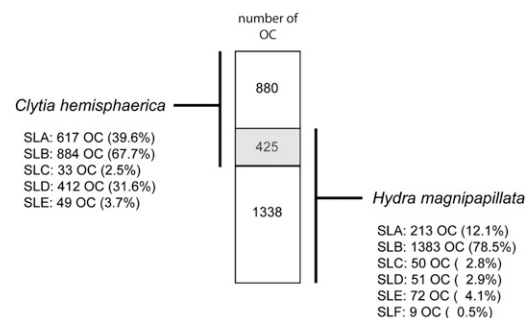
We further showed that the number of Che\_SL groups found spliced to a given mRNA was proportional to the corresponding representation of ESTs in the data set. Thus, OCs *trans*-spliced by one, two, three, and four Che\_SL groups had an average of seven, 21, 34, and 49 corresponding 5' ESTs, respectively (Fig. 3C). Incidentally, these analyses imply that the proportions of mRNAs able to be *trans*-spliced by more than one SL calculated from our OC data

set, as well as the number of distinct SL exons obtained from the whole EST data sets, are almost certainly underestimates.

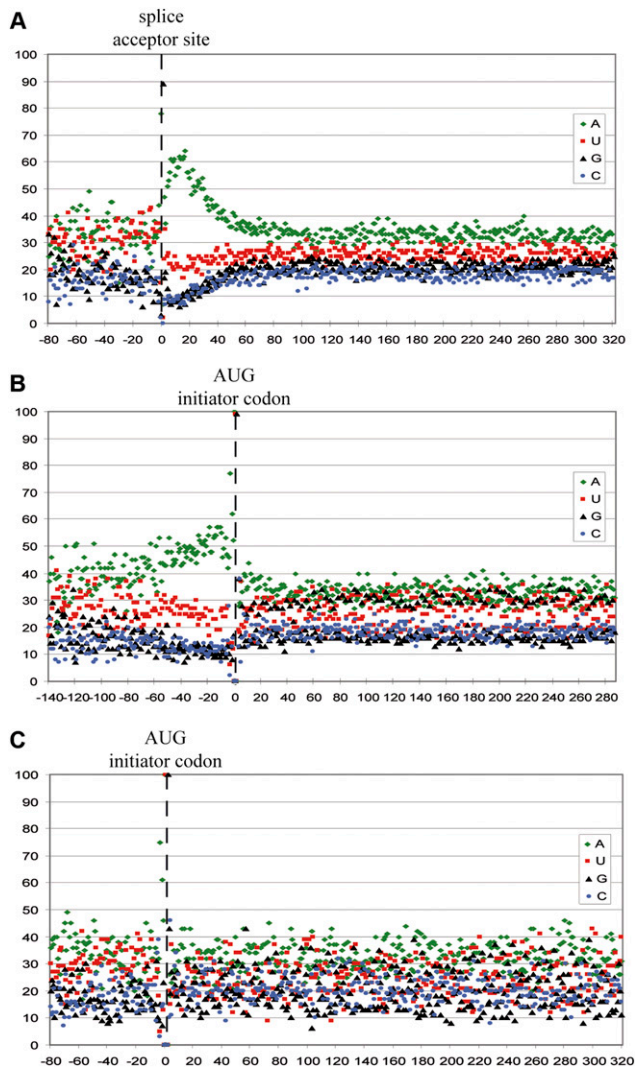
Finally, as in *Pleurobrachia*, we uncovered examples of transcripts *trans*-spliced at alternative sites. These alternative SL *trans*-splicing acceptor sites were found in 9% of OCs *trans*-spliced by at least two SL groups, with the splice acceptor sites always positioned closely together (2–20 nt apart) (Supplemental Data S3).

### Spliced leader *trans*-splicing is favored in adenosine-rich 5' UTRs in hydrozoans

Studies of *C. elegans* polycistronic mRNAs have demonstrated the presence of particular nucleotide contexts favoring SL *trans*-splicing (Graber et al. 2007). In an attempt to detect a similar environment in hydrozoan *trans*-spliced cDNAs, 78 *C. hemisphaerica* cDNA sequences containing outrons (i.e., non-SL 5' termini) were identified by comparing non-*trans*-spliced cDNA sequences with the *trans*-spliced OC data set. We added to this collection 400 further *trans*-spliced cDNAs chosen at random, from which SL sequences were removed and aligned with respect to their splice acceptor site (Fig. 5A). Nucleoside proportions per site along this alignment reveal a marked local enrichment of adenosine just downstream of the splice site (reaching a peak of 65% ~15 bases from the splice acceptor site), whereas no particular enrichment was detected in the outtron. An identical enrichment of adenosine was observed using *H. magnipapillata* *trans*-spliced cDNAs (400 *trans*-spliced cDNAs chosen at random and aligned with respect to their splice acceptor site; data not shown), indicating that this feature is common to these two hydrozoan species. When the *Clytia* cDNA sequence data set was aligned with respect to the AUG translation initiation codon (predicted by GENSCAN), the adenosine enrichment was again detected but was spread more broadly along the 60 bases upstream of the AUG initiation codon (Fig. 5B), indicating that the A-rich area is more likely to be linked to SL *trans*-splicing than to translation initiation. In contrast, no



**FIGURE 4.** Overlap between *Clytia* and *Hydra* *trans*-spliced transcripts. Schematic representation of the orthologous clusters (OCs) data set constructed in this study: Among the 2643 OCs, 425 display *trans*-spliced cDNAs from both species. For each hydrozoan species, percentages of SL groups observed in this OC data set are indicated.



**FIGURE 5.** Analysis of *trans*-spliced and non-*trans*-spliced *C. hemisphaerica* 5' UTRs. (A) Distribution of nucleotide percentages per position along 478 *trans*-spliced cDNAs of *C. hemisphaerica* aligned on the splice acceptor site. (B) Distribution of nucleotide percentages per position along the same data set of *trans*-spliced cDNAs of *C. hemisphaerica* aligned on the predicted ATG initiator codon. (C) Distribution of nucleotide percentages per position along 88 non-*trans*-spliced cDNAs of *C. hemisphaerica* aligned on the predicted ATG initiator codon. In order to avoid stochastic variations of nucleotide percentages in the 5' end of incomplete 5' termini in the assembled ESTs, the positions localized at the 5' end were removed until the presence of at least 50 nt per position is shown in these three graphics.

particular nucleoside enrichment was detected in the 5' UTRs of 88 non-*trans*-spliced cDNAs of *C. hemisphaerica* chosen from conserved and well studied proteins, e.g., ribosomal proteins (Philippe et al. 2009) and developmental regulator genes (Chevalier et al. 2006; Momose et al. 2008; Amiel et al. 2009), except in the Kozac environment (first 3 base pairs [bp] upstream of the AUG initiator codon) (Fig. 5B,C). The random nucleoside distribution in the 5' UTR of non-*trans*-spliced cDNAs argues in favor of a functional link

between the SL *trans*-splicing and the A-rich environment and suggests that certain A-rich 5' UTR compositions are favored for SL *trans*-splicing.

A-rich regions in the 5' UTRs of SL *trans*-spliced transcripts were not detected outside hydrozoans: No particular nucleotide enrichment was detected downstream of the splice site in SL-*trans*-spliced cDNAs identified from the ctenophore *P. pileus* (all 3651 *trans*-spliced cDNAs) and the nematode *Caenorhabditis elegans* (300 cDNAs *trans*-spliced by SL1 plus 300 cDNAs *trans*-spliced by SL2, randomly chosen), or in *trans*-spliced cDNAs identified from ESTs of the urochordate *Oikopleura dioica* (Ganot et al. 2004). It remains possible that more subtle features of the nucleoside context are associated with SL *trans*-splicing in certain mRNAs from these species.

## DISCUSSION

In this study we exploited the availability of large EST data sets to undertake a survey of SL usage in the basal metazoan phyla and related unicellular organisms, and a detailed comparative analysis of SL usage between the hydrozoans *C. hemisphaerica* and *H. magnipapillata*. Our results have shed new light on the origin and evolution of SL *trans*-splicing. They strongly reinforce a scenario in which SL genes have had multiple origins and rapid modification during animal evolution, although of course the possibility of loss of SL usage in one or more evolutionary lineages cannot be ruled out.

Concerning the phylogenetic occurrence of SL usage, our character optimization (Fig. 1) supports its multiple independent origins in ctenophores, hydrozoans, and various bilaterian lineages. This convergence hypothesis is consistent with the high diversity of SL genes in terms of sequences within Eumetazoa and their multiple proposed functions (discussed below). Note that convergence also almost certainly occurred at the wider scale of eukaryotes as unambiguously indicated by the occurrence of SL genes in distantly related eukaryote groups, such as euglenozoans and dinoflagellates. Equivalent conclusions were reached in a parallel study using a very similar approach on EST from a wide range of metazoan species, published while our work was under review (Douris et al. 2009).

An attractive hypothesis to explain multiple evolutionary origins for the SL genes is that they have derived repeatedly from U-rich small nuclear RNAs (snRNAs) of the Sm-class involved in the nuclear spliceosome machinery (for reviews, see Nilsen 2001; Hastings 2005). Like the SL RNAs, these snRNAs (U1, U2, U4, and U5) are characterized by a TMG cap and a U-rich Sm-protein-binding site, and are present in multiple copies per genome. The most likely candidate SL ancestor is the U1snRNA, which is the only snRNA absent from the *trans*-splicing spliceosome, and which can acquire *trans*-splicing ability with just a few nucleotide changes (Bruzik et al. 1988; Bruzik and Steitz

1990; Hannon et al. 1992). Duplications of the U1 snRNA gene followed by just a few mutations would be sufficient to lead to the acquisition of *trans*-splicing, since SL pre-RNAs introduced into cells of non-*trans*-splicing species have been shown to undergo *trans*-splicing (Bruzik and Maniatis 1992).

Like the SL genes, snRNA genes occupy inter-5S gene regions in some species (Drouin and de Sa 1995; Ebel et al. 1999; Ganot et al. 2004; Lidie and van Dolah 2007), encouraging hypotheses of an evolutionary relationship. Furthermore, the presence of pseudogenes in 5S rRNA clusters (Jacq et al. 1977) is indicative of frequent gene duplications in this region of the genome. The majority of SL-positive species examined, however, including *Hydra* (this study), do not show this association (Aksoy et al. 1992; Keller et al. 1992; Liu et al. 1996; Zhang et al. 2009), while the U6 genes positioned between 5S rRNA genes *Clytia* and *Laomedea* are clearly unlikely to have a recent evolutionary relationship with the Sm-binding U-rich snRNA group or the SL genes, being transcribed by a different RNA polymerase (Pol III) and lacking their TMG cap. Co-accumulation of SL, snRNA, and 5S rRNA genes may have been favored during evolution in some species, either independently or as a result of co-regulation (Lidie and van Dolah 2007). Further analysis of gene content in 5S rRNA gene clusters, especially in non-SL *trans*-splicing species, may help to resolve this issue (see Drouin and De Sa 1995). Recent analysis in a selection of metazoans uncovered an association of U1 snRNA and SL genes in the amphipod crustacean *Parhyale*, but not in the ctenophore *Mnemiopsis*, the chaetognath *Spadella*, or the bdelloid rotifer *Adineta*, with no 5S rRNA association detected in any of these species (Douris et al. 2009).

Our comparison of SL *trans*-splicing in two hydrozoan species provides a striking demonstration of very rapid SL evolution. Both *C. hemisphaerica* and *H. magnipapillata* exhibit a high number of different SL exon sequences, divided into five and six SL groups, respectively. In the case of *Hydra* we were able to identify all but one of the identified variants as a distinct gene from genome sequence data, indicating that the SL groups represent gene families. The SL groups have clearly diversified rapidly, as shown by a lack of clear sequence conservation between these two distant hydrozoan species, and by the marked divergence from a common ancestral gene of *H. vulgaris* SLA and *H. magnipapillata* SLA1. Since the rate of SL sequence evolution in these species seems similar to that of the intergenic 5S region, and SL exon vs. intron substitution rates are not significantly different, SL sequences appear essentially unconstrained.

The high diversity and rapid evolution of SL in hydrozoans is very similar to the situation reported in the nematode *Trichinella spirallis* (Pettitt et al. 2008) and in the Chaetognaths (Marletaz et al. 2008), as well as in ctenophores (this study), where multiple SL groups from

*P. pileus* and *M. leidy* EST data were found to be closely related but distinct. In contrast, urochordates (Ganot et al. 2004; Satou et al. 2006) and SL-positive unicellular eukaryotes (Gibson et al. 2000; Zhang et al. 2007), show only one SL exon per species (or one SL group), while in Rhabditida nematodes SL1 and SL2 RNAs show functional dichotomy associated with an atypically high level of conservation of the SL1 sequence between species (Blumenthal 2005; Guiliano and Blaxter 2006). The lower level of SL group diversification in these latter groups may be explained in part by stabilization of SL usage following the acquisition of a function in polycistronic transcript resolution in Rhabditida nematodes and urochordates, or changes in the translational machinery to accommodate the altered 5' cap structure in trypanosomes. More detailed analysis of the diversity and evolution of SL sequences in other lineages such as flatworms, acoels, and Bdelloid rotifers may help to confirm whether rapid and high diversification of SL genes is a general rule, and to uncover any potential link between evolution rates and SL function. Flatworms may provide an interesting case, as the AUG initiator codons of some *trans*-spliced mRNAs are provided by the 3' end of SL sequences, which might be predicted to constrain evolution of the SL sequences (Cheng et al. 2006).

In line with the hypothesis that the acquisition (or retention) of specific functions for SL *trans*-splicing in certain lineages might constrain SL gene evolution, we were unable to find any evidence for distinct functions for the diversified hydrozoan SL groups. Firstly, there was no evidence for preferential *trans*-splicing of a particular set of orthologous mRNAs in both *Clytia* and *Hydra*. This tendency was confirmed for particular proteins or protein classes. For instance, no common *trans*-spliced ribosomal protein mRNAs could be identified in the *C. hemisphaerica* and *H. magnipapillata* EST data sets: rpl7a, rpl10, rps27a, and rps35 are *trans*-spliced in *C. hemisphaerica* ESTs, whereas rpl9, rps2, rps6, and rps19 are *trans*-spliced in *H. magnipapillata* ESTs (among 46 and 73 ribosomal transcripts containing 5' UTRs, respectively). Second, the correspondence in *C. hemisphaerica* between SL group diversity for a given mRNA and the number of 5' ESTs in the data set suggests that any of the SL groups can be added to a given mRNA being processed. Furthermore, no preference of particular SLs to splice given mRNAs within a species was found. On the other hand, the different SL variants were not represented proportionally in the hydrozoan EST data set. This suggests that there may be a preference for certain SL genes to be used for *trans*-splicing, although we cannot completely rule out sampling artifacts. Further analysis of SL genes in hydrozoan genomes should clarify this point. Overall, the apparently indiscriminate use of SL sequences indicates that distinct SL groups have no functional specialization, but may share common function(s) such as 5' UTR sanitization.

While we found no evidence for a functional specialization of SL usage in hydrozoans, we did uncover a marked tendency for SL *trans*-spliced transcripts to contain an unexpected local enrichment of adenosine in the 5' UTR prior to splicing, suggesting that a particular nucleoside environment favors SL *trans*-splicing. In the two hydrozoan data sets, SL addition was found to occur ~15 bp upstream of an A-rich region. No such regions were detected in non-*trans*-spliced mRNAs, indicating that pre-mRNAs may be selected for *trans*-splicing on the basis of 5' UTR composition rather than the coded protein. We were not able to detect any nucleoside enrichment in relation to the splice site in other species, although more subtle motifs may be present. Thus, the correlation between SL *trans*-splicing and an enrichment of adenosine in the 5' UTR appears to be a hydrozoan-specific feature of SL *trans*-splicing.

The overall picture of SL evolution emerging from our analyses together with published studies is of frequent SL gene acquisitions, perhaps following mutations in U-rich snRNAs of the spliceosome, followed by rapid sequence evolution and frequent gene duplications. In some cases, modification of the 5' UTR in some or all mRNAs by SL *trans*-splicing may have had consequences for the machinery regulating translation, stability, or operon processing, and thus have constrained subsequent SL sequence evolution (Hastings 2005). As more EST data become available from a range of phylogenetic lineages, this hypothesis can be further examined. The Hydrozoa, a broad and diverse monophyletic group comprising about 3700 described species (Collins et al. 2006), is a promising group in which to study SL *trans*-splicing. *Hydra* and now *C. hemisphaerica* (Houliston et al. 2010) are well-established experimental organisms, allowing experimental manipulation. Moreover, as demonstrated by 5S rRNA cluster analysis, the close evolutionary scale represented by the *Hydra* genus offers an appropriate framework to catch SL sequence divergence. Finally, the newly available genome sequence of *H. magnipapillata*, soon to be joined by that of *C. hemisphaerica* (sequencing ongoing at the Genoscope), will allow large-scale comparative genomic analysis between these two distant hydrozoan species, for instance to investigate the distribution and evolution of operons (Guiliano and Blaxter 2006; Satou et al. 2006).

## MATERIALS AND METHODS

### *Clytia* and *Pleurobrachia* ESTs

EST sequencing for *P. pileus* (~30,000) and *C. hemisphaerica* (~90,000) was performed at the Genoscope from a mixture of normalized and nonnormalized cDNA libraries, constructed by Open Biosystems (through BioCat) and Express Genomics from microgram quantities of total RNA extracted from adult (Pp) or mixed embryonic, larval, and adult (Ch) stages (Chevalier et al. 2006; Philippe et al. 2009). All starting material was obtained from

Villefranche-sur-Mer, with the *C. hemisphaerica* derived uniquely from three cultured strains (X, Y, and Z). These EST sequences are available in dbEST/GenBank (<http://www.ncbi.nlm.nih.gov/dbEST/>).

### Detection of SL sequences in EST data sets

To recover SL exon sequences, the following steps were performed independently on each EST data set. The method was validated by recovering known SL exons from *C. elegans* ESTs and from *H. vulgaris* ESTs. Cleaned ESTs with vector sequences removed were assembled into contigs using Phrap software. The assemblies were then searched for common sequences of 12 nt at the termini, and all elements present at least three times were aligned manually to reconstruct putative SL sequences. The putative SL sequences were then used to search by nucleotide BLAST the original assembled cDNA data set, and those found identically in at least five 5' ends of contigs were defined as SL variants. SL variants displaying three or less nucleotide changes then were considered to form a SL group.

### Amplification of 5S rRNA gene repeats in hydrozoans

Genomic DNA was extracted from *C. hemisphaerica* male medusae by standard methods. *L. calceolifera* genomic DNA was provided by Peter Schuchert (Muséum d'histoire naturelle de Genève) and Lucas Leclère (Sars Institute). Amplification of inter-5S rRNA regions was performed using primers and PCR cycles as defined in Stover and Steele (2001), and inserted into pGEMt plasmid for sequencing.

### Construction of hydrozoan *trans*-spliced orthologous clusters

Hydrozoan orthology clusters were compiled using the *N. vectensis* proteome as reference (Putnam et al. 2007), downloaded from the JGI website (<http://www.jgi.doe.gov/>). Each *N. vectensis* protein defines one OC. SL *trans*-spliced assembled cDNAs from *C. hemisphaerica* and *H. magnipapillata* ESTs were compared by BLAST against these protein sequences and were assigned to the OC corresponding to their best hit using a threshold value of 1e-10. For all OCs showing multiple SLs and for a selection of 250 among those with single SLs, we also retrieved from the original ESTs all orthologous non-SL *trans*-spliced transcripts for analysis of splice context. For each OC, cDNAs were aligned using MUSCLE (Edgar 2004), independently for each hydrozoan species, and alignments obtained were carefully checked by eye in order to detect the presence of different transcripts. Rare cases (<5%) in which at least two distinct transcripts were assigned to a single OC for one or both species were discarded to avoid complex multigenic families. The orthology of *trans*-spliced cDNAs from *Clytia* and *Hydra* was confirmed by successful phylogenetic analysis for 20 OCs chosen randomly among the 425 OCs displaying *trans*-spliced cDNAs from both species (data not shown).

### Accession numbers

The GenBank (<http://www.ncbi.nlm.nih.gov/Genbank>) accession numbers for the 5S rRNA gene sequences identified in this study are GU126459 (*C. hemisphaerica*) and GU126460 (*L. calceolifera*).

## SUPPLEMENTAL MATERIAL

Supplemental material can be found at <http://www.rnajournal.org>.

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