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Authors
Tarrío, R
Ayala, FJ
Rodríguez-Trelles, F

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Alternative splicing: A missing piece in the puzzle of intron gain

Rosa Tarrio*,†, Francisco J. Ayala†‡, and Francisco Rodríguez-Trelles*†‡

*Grupo de Medicina Xenómica–Centro de Investigación Biomédica en Red de Enfermedades Raras, Hospital Clínico Universitario, Universidad de Santiago de Compostela, 15706 Santiago, Spain; and †Department of Ecology and Evolutionary Biology, University of California, Irvine, CA 92697-2525

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Spliceosomal introns, a hallmark of eukaryotic gene organization, were an unexpected discovery. After three decades, crucial issues such as when and how introns first appeared in evolution remain unsettled. An issue yet to be answered is how intron positions arise de novo. Phylogenetic investigations concur that intron positions continue to emerge, at least in some lineages. Yet genomic scans for the sources of introns occupying new positions have been fruitless.

Two alternative solutions to this paradox are: (i) formation of new intron positions halted before the recent past and (ii) it continues to occur, but through processes different from those generally assumed. One process generally dismissed is intron sliding—the relocation of a preexisting intron over short distances—because of supposed deleterious effects. The puzzle of intron gain arises owing to a pervasive operational definition of introns, which sees them as demarcated segments of the genome separated from the neighboring nonintronic DNA by unmovable limits. Intron homology is defined as position homology. Recent studies of pre-mRNA processing indicate that this assumption needs to be revised. We incorporate recent advances on the evolutionarily frequent process of alternative splicing, which exons of primary transcripts are spliced in different patterns, into a new model of intron sliding that accounts for the diversity of intron positions. We posit that intron positional diversity is driven by two overlapping processes: (i) background process of continuous relocation of preexisting introns by sliding and (ii) sprouts of extensive gain/loss of new intron sequences.

Background

Intron Positions Arise de Novo in Evolution. Approaches to the evolution of intron positions have become increasingly sophisticated since the early comparisons of GenBank data (1). Yet the prevalence with which new intron positions arise in evolution continues to be debated (2–5). At the root of the controversy are differences in methodological postulates, phylogenetic sampling scopes, and criteria for deciding intron positions.

Ancestral intron positions are inferred from a matrix of intron presence/absence built by projecting present positions onto automated multiple sequence alignments of genome scale sets of orthologous proteins. Rogozin et al. (6) compiled 684 clusters of orthologous genes (KOGs) from eight model eukaryotes, including one vertebrate (human), two arthropods (Drosophila melanogaster and Anopheles gambiae), one nematode (Caenorhabditis elegans), two fungi (Saccharomyces cerevisiae and Schizosaccharomyces pombe), one plant (Arabidopsis thaliana), and one protist (Plasmodium falciparum). The resulting 16,577 unique intron positions were condensed into 7,236 (~43%) by retaining only those located within well conserved tracts of alignment. The full and conserved matrices were analyzed by Dollo parsimony (6). The conserved matrix was subsequently reanalyzed by other authors. Roy and Gilbert (7) devised a local maximum-likelihood (ML) approach that corrects for the known bias of Dollo parsimony toward the overestimation of intron gain at peripheral branches, owing to a failure to detect intron losses that are not directly observed. However, when the number of target sites (i.e., observed plus unobserved intron positions) is taken into account explicitly in ML simultaneous comparison of all species (8–10), the numbers of ancestral intron positions are fewer than those obtained previously (7). The reason could be that the method of ref. 7 does not allow for homoplastic gains (i.e., introns arising more than once at the same homologous position) (8, 9, 11), but it also could be that homoplastic gains are overestimated by ML methods (e.g., due to sparseness of phylogenetic sampling). Homoplastic gains seem to have been extremely overestimated by Qiu et al. (12), who claim that the vast majority of intron positions are new apparently because, in their Bayesian analysis of 10 gene families, the number of target sites is bounded to be equal to the number of observed intron positions (8, 9).

The dataset shown previously (6) has been expanded from 8 to 18 eukaryotic species using a new criterion to determine intron positional homology (10). The 10 added species split long branches of the tree near the tips. The result is a 30% reduction of KOGs (from 684 to 483), but not of intron positions in the matrix, which increases by 10% (from 7,236 to 8,044), almost twice the value that obtains (4,136) by extrapolating from the corresponding numbers in the conserved dataset of ref. 6. A factor contributing to the increase in intron positions may be that ref. 10 rewards matching of intron positions to help align the amino acids, which relaxes the minimum of protein conservation required for identifying intron positions. The ref. 6 dataset also has been expanded by ref. 11 by adding 11 species, 6 of which are not included in ref. 10. Previous models allow for variation of the rates of intron gain and loss, among either species, 6 of which are not included in ref. 10. Previous models allow for variation of the rates of intron gain and loss, among either lineages (7–10) or genes (12), and the Carmel et al. (11) model accommodates both, plus rate variation among sites within a gene, thus avoiding the difficulty of having to estimate the number of target sites separately (8–10). Five of the 11 new species involve intron positions in deuterostomia. The other five species (except for Oryza sativa, which is closely related to Arabidopsis) belong to new long peripheral branches. The increase in the number of species results in a 40% reduction of KOGs (from 684–391) and a 20% reduction in the number of analyzed intron positions (from 7,236 to 5,755).

Most of the studies cited above agree that the last eukaryotic common ancestor (LECA) had a high intron density. A fraction (10–40%) (1, 3, 6) of the ancestral introns has persisted to the present time, although the degree of ancestral intron retention varies among species owing to vast differences in rates of intron loss. But the inferred and/or observed intron positions at many nodes cannot be explained without also invoking differences in rates of gain. Patterns of gain appear to be due to episodic bursts super-

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*To whom correspondence may be addressed. E-mail: fjayala@uci.edu or ftreilles@usc.es.

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imposed on much lower background rates. Except for those spurs, intron losses dominate over intron gains in most lineages (11). Apparently, both gain and loss rates have decreased during the last tens to hundreds of million years, but at a rate decrease much greater for intron gain than for intron loss (10, 11).

Mechanisms for the Origin of Novel Intron Positions. There are at least three global mechanisms for the de novo origin of intron positions: (i) transposition, which would include duplication of preexisting introns; (ii) insertion of intron-like transposons; and (iii) tandem duplication of exon sequences that happen to include splice sites (4, 13). These mechanisms assume that (i) every new intron position originates from a “formative” intron, (ii) formative introns derive from intron donors elsewhere in the genome (including introns, transposons, and exons), and (iii) formation of a novel intron position is instantaneous. Formative introns are at first identical to their donors and are expected to remain detectably similar for millions of years. A straightforward approach to show that intron positions arise by any of the proposed mechanisms is finding the donors of formative introns, which should not be difficult provided recent intron positions arise in sufficient numbers.

Results

Recent Rates of Origination of New Intron Positions. Rates of intron gain are inferred to have strongly declined during the last tens to hundreds of million years (6, 8–11). Table 1 (“Long-term”) shows the rates of intron gain for long peripheral branches, not appropriate for evaluating recent gains, but given for comparison. The rates tend to decrease as the complexity of the evolutionary model increases. The lowest values are attained by allowing for gain rate variation among lineages, genes, or sites (11). The approach begins by identifying discordant intron positions between closely related homologs. The discordant positions are then compared to an outgroup. The discordances that match an intron in the outgroup are attributed to intron loss; otherwise they are attributed to intron gain. Roy et al. (14) found no evidence of intron gain from 1,560 human-mouse and 360 mouse-rat orthologs (using the fish *Fugu* and human as outgroups, respectively).

No case of gain was reported in a mapping of annotated intron–exon boundaries of either 17,242 human or 16,068 mouse genes in alignments of human, mouse, rat, and dog genomic sequences (17) (this result appears to be at variance with that obtained by ref. 25, which reported many novel introns in humans, although the new intron-containing genes are either unannotated or in copy-number variant regions). *D. melanogaster* (subgenus *Sophophora*) is inferred to have gained ∼0.45 introns/gene/By during the ∼40 My elapsed since it split from the *Drosophila* subgenus (18). Table 2 gives parsimony estimates of intron gain from closely related species/lineages.

The higher ML rates of recent intron gain, compared with those obtained with parsimony, cannot be accounted for by systematic differences in calibration dates between the two optimality criteria. Under a range of models, parsimony is an ML estimator, but not for the model that allows multiple changes (gains or losses) at a position (26). Intron gain/loss has only two alternative states and, thus, is more vulnerable to homoplasy. Homoplastic gains represent 5–20% of shared intron positions (8, 9, 27, 28). Although the potential for homoplastic gain decreases with the divergence in the sample, closely related sequences are prone to it by virtue of their high similarity (provided gains do not occur at random) (29). Studies of closely related species that use distantly related outgroups (e.g., 15, 17, 21, 22) have enhanced likelihood of parallel gain. However, both the ML and parsimony estimates would be downwardly biased if newly gained intron positions tend to be excluded by data filtering.

To avoid database errors in intron–exon boundaries and annotation, analyses of intron gain are typically confined to positions in windows of protein alignment that are highly conserved and often do not contain gaps (6, 8–11, 14–18, 20, 21, 24). In addition, slight
Table 2. Parsimony estimates of recent rates of intron gain (per gene per 10^9 years) for some better studied lineages

<table>
<thead>
<tr>
<th>Ref(s)</th>
<th>Gene set</th>
<th>Rate (intron/gene/By)</th>
<th>Lineage* (tree; calibration time in My)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15, 22</td>
<td>16,590h</td>
<td>0.0034–0.0127</td>
<td>C. elegans (((((Cb,Cr,Csp4),(Cg,Em),(Hs),(Sp)); 100</td>
</tr>
<tr>
<td>24</td>
<td>4,690h</td>
<td>0.0023</td>
<td>C. neoformans (((Cn,Cgl),(CgR,CgW)); 37</td>
</tr>
<tr>
<td>16</td>
<td>1,447h</td>
<td>0.66</td>
<td>Three ascomycete fungi ([(Mg No),(Fl,An); 630</td>
</tr>
<tr>
<td>19, 23</td>
<td>2,563p</td>
<td>0.30–0.90</td>
<td>A. thaliana (((At1,At2),Le),(Ox); 20–60</td>
</tr>
<tr>
<td>20</td>
<td>3,101p</td>
<td>0.15</td>
<td>O. sativa (((Ox1,Ox2),At2); 70</td>
</tr>
<tr>
<td>21, 26</td>
<td>3,479p</td>
<td>0.0014–0.0115</td>
<td>P. falciparum (((((Pv,Pf),(Pf),(Pf),(Ta,Tp)); 100</td>
</tr>
</tbody>
</table>

The number next to each tree is the duration (My; from references in the leftmost column) of the branch over which rates are calculated (underlined). h, homolog; p, paralog.

*Species not given in Table 1 are Cr and Csp4, Caenorhabditis remanei and sp. d., respectively; Bm, Brugia malayi; Sp, Schizosaccharomyces pombe; Cnn and Cng, Cryptococcus neoformans var. neoformans and C. neoformans var. grubii, respectively; Cgl and CgW, Cryptococcus gattii strains R265 and WM276, respectively; Mg, Magnaporthe grisea; Nc, Neurospora crassa; Fg, Fusarium graminearum; An, Aspergillus nidulans; At1 and At2, Arabidopsis thaliana duplicates 1 and 2, respectively; Le, Lycopersicon esculentum; Os1 and Os2, Oryza sativa duplicates 1 and 2, respectively; Pv, Py, Pf, and Pg, Plasmodium knowlesi, P. vivax, P. yoelli, P. falciparum, and P. gallinaeum, respectively; Ta and Tp, Theileria annulata and T. parva, respectively.

discordances (<6 nt) are either excluded (14, 16, 21) or treated as orthologous (6, 8–11). Consequently, conclusions about the incidence of intron gain are based on small subsets (≤50%) of the positional discordances in the unfiltered data (6, 8–11, 14, 17, 20, 30). But the subsets will be impoverished in gained intron positions if the mechanisms that create new positions can cause indels and/or changes in the amino acid composition of flanking exons (see below).

An indication that filtering methods may underestimate the rates of intron gain comes from ref. 6, where the ratio between gains and losses in the unfiltered dataset (5,377/14,341 = 2.67) is ~50% greater than in the conserved dataset (3,306/5,951 = 1.80). The proportion of counted gains at peripheral branches also is greater for the full dataset (50% vs. 40% for the full and conserved datasets respectively, excluding deep-branching fungi, Arabidopsis, and Plasmodium). It seems reasonable to assume that the rates of recent intron gain are somewhere between the ML and the parsimony estimates. Some parsimony rates would still imply substantial gains of introns in the recent past when extrapolated to a genomic gene number scale. In particular, the number of introns gained/My would be ~6.1 in the lineage of D. melanogaster (assuming 13,600 genes), between 7.6 and 22.9 in A. thaliana (25, 500), 7.5 in rice (50,000), and 6.6 in Ascomycete fungi (10,000).

The Puzzle of the Origin of Novel Intron Positions. Genome scans carried out in human, D. melanogaster, C. elegans, and A. thaliana could not detect a single case of homologous introns in nonhomologous genes (31). Subsequent studies in Drosophila, Caenorhabditis, and rice searching specifically for donors of introns supposed to be novel because of restricted phylogenetic distribution (15, 18, 20) also have been fruitless (7, 18, 22, 23). This may not be surprising for humans and nematode if one assumes that the rates of recent intron gain are between 7.6 and 22.9 in D. melanogaster, a 100-bp-long formative intron between 1.5 and 10.0 in C. elegans, and rice even if we assume a rate of 1.0% for the recent past when extrapolated to a genomic gene number scale. The genome became identified with how it is effected through splicing of a particular mRNA. This formal identification fostered a categorization of introns as precisely demarcated segments of the DNA level, was implicit in the experimental observations that led to the discovery of introns (36). The fact that the AS products were of the conceptualization of introns as definite DNA segments. Yet seeking to identify mechanisms for the rapid evolution of protein-coding sequences (34, 37–39), records were cited of so-called cryptic donor/acceptor splice sites (40, 41), and speculations were advanced that splicing-altering mutations could cause extensions/contractions of exons at intron junctions.

Intron Sliding in a World Lacking Alternative Splicing (AS). AS, and the notion that there is not necessarily a one-to-one correspondence between intronic DNA and splicing products at the RNA/protein level, was implicit in the experimental observations that led to the discovery of introns (36). The fact that the AS products were of the casette type (i.e., exons that are alternatively included/skipped from the mature transcript) did not question the emerging conceptualization of introns as definite DNA segments. Yet seeking to identify mechanisms for the rapid evolution of protein-coding sequences (34, 37–39), records were cited of so-called cryptic donor/acceptor splice sites (40, 41), and speculations were advanced that splicing-altering mutations could cause extensions/contractions of exons at intron junctions.

The discovery of increasing examples of lineage-specific introns (e.g., 42–44) launched the debate on the origins of new intron positions. The hypothesis of intron sliding (IS), also named “intron drift,” “intron migration,” or “intron slippage,” holds that new intron positions arise by the relocation of preexisting introns (45, 46). Relocation events would take place through the reassignment of an intron’s donor and acceptor splice junctions to nearby positions, both offset in the same direction by the same distance (47, 48). But owing to its likely stepwise mechanism (see below), IS may
An AS-Driven Model of IS. At the time that the hypothesis of IS was launched, AS poorly annotated genes with respect to AS (1, 6, 10, 11, 21, 22, 47), and including positional homology of introns. OneMoreover, phylogenetic approaches, which provide the evidence standing formulations neglect that AS can facilitate the process. IS would increase the diversity of intron positions without increasing the number of introns. Hence, IS would not be a valid explanation for introns in intron-bearing genes that were previously intronless, such as processed pseudogenes (although initial intron positions may slide later).

Interest in IS models diminished on the belief that IS could not be a frequent phenomenon (4, 6, 47, 48). Under the notion of introns as fixed genomic segments, IS is perceived as uncommon because it calls for the simultaneous occurrence of two mutations. Other paths, by a series of two or more short-range extension/contraction events of intron–exon boundaries, were deemed likely to be deleterious at the protein level (47, 48). Such events would be feasible when the aberrant mRNAs contained premature stop codons that could be targeted by nonsense mediated decay (NMD) (49). Provided the locus is haplosufficient, degradation of the transcript would turn out the mutant allele completely recessive, (49). Provided the locus is haplosufficient, degradation of the transcript would turn out the mutant allele completely recessive, (49).

IS is thought to exhibit low potential for intron relocation because standing formulations neglect that AS can facilitate the process. Moreover, phylogenetic approaches, which provide the evidence for the incidence of IS, have overlooked AS as a fundamental consideration in deciding the positional homology of introns. One reason for this neglect is that homologous intron positions have largely been established by extrapolation from unannotated or poorly annotated genes with respect to AS (1, 6, 10, 11, 21, 22, 47, 48, 50, 51). At the time that the hypothesis of IS was launched, AS was still thought to be a minor processing pathway (52).

An AS-Driven Model of IS. New splice sites can arise by point mutation because donor and acceptor splice sites are short and imprecise (53). Any gene region likely includes many more donor and acceptor splice sites than those implied by the exon junctions of mature transcript molecules (54–56). There is not a one-to-one correspondence between donor and acceptor splice sites. One donor may pair with more than one of several acceptors and the other way around, giving rise to a profile of AS products or transcript isoforms, which can differ in the exons they contain, but also in the location of exon junctions (56, 57). Alternative mRNA isoforms evince that fixed intron locations are not suitable for determining positional homology at the genome (DNA) level.

AS has been reported in animals, fungi, plants, and various protists and was probably present in the intron-rich LECA (58). Many AS events, especially those involving weak splice sites, are idiosyncratic across species (38, 59–62). Most AS events can be classified into four basic patterns, including exon skipping, alternative 3’ and 5’ splice site selection, and intron retention. The patterns required for IS, namely, alternative 3’ and 5’ splice site selection, are the most or the second most prevalent type of AS event, accounting for at least one third of all AS events in invertebrates, vertebrates, and Arabidopsis (55–57). A typical human gene may yield 2.53 splicing isoforms translatable to protein (63). Such a diversity of mRNAs and proteins may, in part, be redundant and carry out new functions and may not be “visible” to natural selection (38, 39, 63, 64). However, a substantial fraction will involve changes unlikely to be tolerated (63, 65–67).

Donor-acceptor splice pairs can be strong or weak variants according to frequency of use. Strong splice pairs yield major isoforms, present in >50% of the transcripts of an allele, whereas weak splice pairs yield minor isoforms, which are a small fraction of the normally spliced, mature mRNA (38, 39). Differential production/processing of transcript isoforms may be at the core of organismal robustness to the diversity of AS products (38, 39, 63). It has been proposed that newly arising, potentially deleterious AS products convey only weak splice signals and, hence, are minor isoforms (38, 60). Because of their low abundance, minor isoforms would not often have a major impact on physiology; thereby, they would evolve relatively unconstrained, provided the major fraction of transcripts upholds the gene’s function (38). So-called “tunneling” of aberrant AS forms enhances their retention in a population, which increases the likelihood of compensatory mutations to a restored or novel function if they happen to be disclosed to selection (38, 64, 68). Unlike the standing model of IS via NMD (49), in IS via AS, a second genomic copy of the gene would not be required to maintain the original function because AS would furnish internal paralogs of the gene. This hypothesis is supported by a study showing that (i) minor-form AS relaxes selection pressure against premature termination codons (PTCs) that are likely targets of NMD (to the same degree as having two copies of the gene), and (ii) the combined effects of AS and diploidy yield a >9-fold increase in tolerance for PTCs (69). By enhancing the rate of compensatory mutations, AS expand the potential paths to IS over those under NMD. The threshold of approximately four codons above which IS is considered to be unviable (47, 70) is most likely an underestimate.

The relative use of a given donor–acceptor splice pair depends on the interactions between trans-acting factors and the splicing code. The splicing code is made up of an extensive and complex array of cis-acting elements featuring two layers of information. The first layer comprises the splice site sequences that define potential intron–exon junctions on the target pre-mRNA. The second layer consists of splicing enhancers and silencers distributed all over the introns and exons of the target pre-mRNA. This second informational layer determines which and with what frequency splice sites of the first layer will become targets of the trans-acting factors (71).

The interactions between trans- and cis-acting splicing elements are highly context-dependent. Every site of a pre-mRNA molecule can potentially influence the production of a transcript isoform (55, 56), which implies that there is an extensive genomic target for mutations that can affect AS profiles. This conclusion is supported by the large and growing number of inherited human diseases found to be caused by AS-altering mutations (56, 71, 72). Likely, those mutations represent only extreme cases of an abundant class of genetic polymorphisms that generate quantitative variation in the ratios of isoforms among individuals (73–75). The mutations responsible for this variation may spread and become fixed or lost under the forces of population genetics, just like any genetic variant. Minor splice isoforms would evolve into major isoforms, replacing preexisting predominant gene products, which would then become minor isoforms and be lost over time. The discovery of ancient human pseudogenes, originated by reverse transcription of AS products not presently expressed by the parent gene (76), suggests that the strength of a splice site is dynamic during evolution. This idea is further supported by observations that AS profiles tend to diverge rapidly after gene duplication (77) or speciation events (61, 78). If a preexisting major isoform is superseded by another isoform bearing expansions/contractions of exon limits or slid exon junctions, the replacement would cause a change in the distribution of intron positions of the gene (see Fig. 1).

De Novo Origin of Intron Positions: Intron Sliding Versus Intron Gain. The arguments given suggest that AS could provide a major avenue for the occurrence of IS, one that may have been seriously underestimated as a source of intron positional diversity. A reason that IS has been disfavored over gain of new introns in accounts of intron positional diversity is the assumption that IS must involve large deleterious effects (47, 48). However, increasing understanding of
the complexity of splicing codes suggests otherwise. Optimal splicing codes must require time to evolve (79). It seems unlikely that a de novo intron-formation event, regardless of whether it derives from another intron, a transposon, or an exon donor, can lead to an immediately efficiently spliced product. If splicing of a formative intron is inefficient, then the unspliced, intron-retaining, and, hence, unlikely to be functional transcript will set off as a major isoform, hence exposed to negative selection. Thus, the creation of intron positions from new introns may have larger fitness costs than IS of preexisting introns over short distances because the latter would take place through the readjustment of preexisting splicing codes via changes in minor isoforms.

IS events are not expected to occur instantaneously. After the emergence of a novel donor/acceptor splice site, millions of years might be necessary until the fixation of the mutation(s), as well as the occurrence of changes in splicing code allowing for the replacement of preexisting major isoforms. At the process completion, little may be left of the original intron sequence. IS events may be more easily detected by retracing phylogenetically the AS events that led to the intron relocation than by interspecific comparison of intron sequences. Comparing closely related genomes, such as those of 12 Drosophila species (80), may help identify such footprints. The persistence of alternative isoforms for long periods of evolutionary time would provide a natural path to parallel gain of intron positions if, after a duplication/speciation event, the same isoform replacement takes place in more than one descendant lineage.

IS may help explain the observed preference of introns to be located at mAG/Gt contexts (where “m” can be A or C, uppercase letters indicate a stronger preference, and “i” denotes the placement of the intron), termed “protopsplice sites.” In actin genes, elimination of normal splice sites in a gene triggers AS of the mutant transcripts via use of cryptic splice sites, which happen to coincide in location with functional splice sites in other orthologs of that gene (51). Newly activated donor (GT) and acceptor (AG) splice sites exhibit a bias to be flanked, respectively, by AG and GT dinucleotides at the exonic side. However, IS may be instrumental to understand reported correlations between intron positions and structural/functional features of the encoded proteins if less harmful AS events have a greater associated likelihood of compensatory mutation.

Gain of an intron position by IS implies loss of the previous position of that intron. IS should generate a strong and positive correlation between the rates of intron gain and loss. Such a correlation has been reported in a recent ML reconstruction of intron evolution in 19 model eukaryote species (11). The study partitioned intron evolution into three modes: balanced mode, characterized by proportional gain and loss rates, and elevated loss or gain modes. Rates of gain and loss were found to be positively correlated only for the balanced mode, as expected of IS, which cannot either create or remove intron sequences. These results suggest that the diversity of intron positions may be dominated by two main effects: a background effect due to the continuous relocation of introns by IS, superimposed by episodes of active addition/removal of new intron sequences by intron gain/loss mechanisms. In this respect, it is important to pinpoint that AS may contribute to the evolution of the diversity of intron positions not only as a catalyst of IS, but also as a potentially powerful mechanism of intron gain. Indeed, a large-scale analysis of the role of AS in exon creation and loss during vertebrate evolution (81, 82) found that new alternative exons set off as minor splice forms in most cases. These minor splice forms originate via mutations that introduce new splice sites inside preexisting intron sequences. Exonization of an intron’s partial sequence effectively splits the original intron sequence in two, thus increasing the initial intron number by one. Because this process creates new introns from separate parts of preexisting introns, it cannot be identified by intragenomic similarity searches. AS-driven exonization of intron partial sequences complements our current knowledge of molecular mechanisms of intron gain. The efficiency of this mechanism should increase with intron length.

If IS is an important determinant of the diversity of intron positions, then it might be expected that the rate of intron position evolution would be positively correlated with the rate of sequence evolution. The two types of evolution depend on the same set of mutations (i.e., point mutations that by changing the sequence would influence the rates of splicing code evolution). The issue has not been investigated in depth, but there are some indications that such a correlation may occur. The sea anemone Nematostella vectensis, the marine annelid Platynereis dumerilii, and humans evolve more slowly than Caenorhabditis and Drosophila at the sequence level. Apparently, they also share larger numbers of ancestral intron positions (83, 84), although anemones and annelids are more distantly related to humans than nematodes and flies. Sverdlov et al. (28) reported a shortage of conserved intron positions in ancient eukaryotic paralogs compared with the higher rate of conservation of intron positions in more recent paralogs. This finding would be consistent with an effect of IS, taking into account that widespread AS appears to be an ancient feature (58), as well as the tendency of AS patterns to diverge after duplication (69, 77).

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