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Mystery of Intron Gain

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For nearly 15 years, it has been widely believed that many introns were recently acquired by the genes of multicellular organisms. However, the mechanism of acquisition has yet to be described for a single animal intron. Here, we report a large-scale computational analysis of the human, *Drosophila melanogaster*, *Caenorhabditis elegans*, and *Arabidopsis thaliana* genomes. We divided 147,796 human intron sequences into batches of similar lengths and aligned them with each other. Different types of homologies between introns were found, but none showed evidence of simple intron transposition. Also, 106,902 plant, 39,624 *Drosophila*, and 6021 *C. elegans* introns were examined. No single case of homologous introns in nonhomologous genes was detected. Thus, we found no example of transposition of introns in the last 50 million years in humans, in 3 million years in *Drosophila* and *C. elegans*, or in 5 million years in *Arabidopsis*. Either new introns do not arise via transposition of other introns or intron transposition must have occurred so early in evolution that all traces of homology have been lost.

[A. Smit and P. Green kindly provided computer programs.]

When introns were first discovered in 1977, an immediate question was where do they come from? Very early in the debate, Crick and others suggested that new introns might arise as transposons that either come equipped with or quickly acquire signals sufficient for splicing (Crick 1979). This idea was reinforced by the finding by Dibb and Newman (1989) that introns tend to arise at sites with a consensus sequence of (C/A)AG|R, which they interpreted in terms of sequence-specific targeting of some not-yet-characterized insertion machinery. However, small-scale comparisons of intron sequences have turned up no evident transposition events, and there is to date a sole convincing demonstration of the creation of an intron from a transposable element, in plants (Giroux et al. 1994).

Yet, it is becoming increasingly clear that some introns have arisen very recently. There is a growing collection of introns found at positions in one species at which closely related species have no intron (Palmer and Logsdon Jr. 1991; Logsdon Jr. et al. 1995; Rzhetsky et al. 1997; Frugoli et al. 1998; Gotoh 1998; Tarrío et al. 1998; Venkatesh et al. 1999; for an insightful review, see Logsdon Jr. et al. 1998; Lynch and Richardson 2002). Over a longer time scale, only 14% of animal intron positions match with the intron positions of homologous plant genes, indicating that, if gain is a primary mechanism of intron discordance, 60%–80% percent of contemporary animal introns were acquired after the evolutionary divergence of animals and plants (Fedorov et al. 2002).

Such new introns should not be hard to find. For instance, the human genome has 31,000 intron-containing genes with 160,000 introns. If 60% of these introns were acquired since the animal–plant divergence then there have been an average of 20–50 cases of intron insertion per million years of human evolution. This estimation is also in accordance with the intron turnover rates of ~0.65–0.70 per billion years for worms and flies (Moriyama et al. 1998; Kent and Zahler 2000; Lynch and Richardson 2002). Regardless of the relatively rapid rate of intron sequence drift, recently acquired introns should still have sequences that betray their origins. Therefore, if introns are, in-

deed, mobile elements, one should be able to detect many transposition events by a comparison of all introns in a genome. Conversely, a failure to find similarities between introns would strongly imply that intron transposition is not a common event in recent evolution.

We obtained intron sequences for all known human genes. Intron sequences were aligned against each other to search for homology. Each intron pair showing significant homology and belonging to nonhomologous genes was then analyzed individually. Among all these inspected cases of intron pairs with strong homology, none showed a relationship indicative of intron transfer. The analogous analysis of *Drosophila melanogaster*, *Arabidopsis thaliana*, and *Caenorhabditis elegans* introns was similarly unable to detect a single recent transposition of an intron from one gene into another. We conclude that either introns do not arise by transposition or that the rate of intron gain has been highly nonuniform through evolution.

RESULTS

Comparison of Human Introns

We divided 147,796 human introns into batches according to their lengths with introns of lengths 1–50 nt constituting batch 0, 51–100 nt in batch 1, and so on. A second set was generated with batch 0 containing 26–75 nt, batch 1 containing 76–125 nt, and so on. This ensures that each pair of introns whose lengths differ by 25 or less is compared and that most pairs with differences <50 nt are compared. Figure 1 shows the sizes of these batches of introns. Within each batch, every pair of introns was compared. The results of these comparisons are summarized in Table 1. Of 210 million pairs of introns that we compared, 13,435 gave BLAST scores of 100 bits or better. Of these, the vast majority were comparisons between introns from homologous genes, indicating that the homologies were not caused by intron transposition but by duplication of the entire original gene. Excluding these left 1368 intron pairs. We next excluded those pairs that came from the same contig. The reason for this exclusion is as follows. There are tens of thousands of short duplications in the human genome. The majority of these DNA fragments lie in the vicinity of their original copies, on the same chromosome. We found the highest level of falsely computer-predicted introns in

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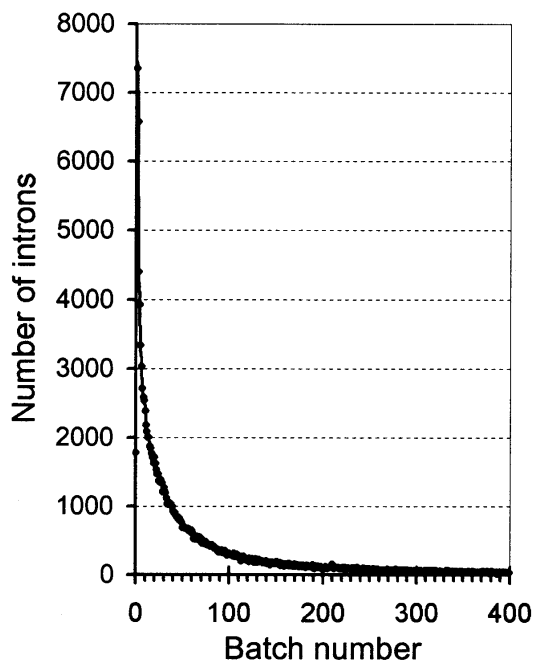


Figure 1 The number of introns in the batches from set one. Batch n consists of all human introns with the lengths between $(50n + 1)$ and $(50n + 50)$ nt. The distributions for batches from set two are similar.

these DNA duplication fragments, mostly arising from dubious predicted genes with no amino acid sequence homology to any known gene. The disposal of introns from the same contig significantly simplifies further analysis but leaves a chance of missing special cases of intron transposition. Hypothetically, insertions of new introns could occur within spatially restricted genomic domains. However, our recent large-scale analysis of human and mouse introns (Roy et al. 2003) makes this conjecture highly unlikely. Removing these closely located intron pairs left 118 pairs of introns in which the members came from non-homologous genes from different contigs.

Figure 2 shows one of these 118 BLAST alignments, between part of the first intron of a computer-predicted gene from Chromosome Y encoding the hypothetical protein XP_067115 and part of the tenth intron of a computer-predicted gene from Chromosome Y encoding the hypothetical protein similar to transcript Y7. These introns are 475 and 469 nt long, respectively, with a 135-nt region of 87% identity (Fig. 2), yielding a BLAST score of 113 bits (e -value 10^{-26}). When this 135-nt region was used to query all of GenBank, it was found that the sequence has 83% identity to testis-specific transcript, *Y-linked 2* (*TTY2*) mRNA, and also is a part of boundary 2 of the *TTY*-like array, a Y-chromosome landmark. Therefore, these two introns are homologous because of the presence of a low-copy Y-chromosome DNA-repeat.

Several other homologous intron pairs had similar explanations. For example, intron 3 of a gene similar to cytochrome oxidase I from Chromosome 9 (16103 EID identifier) and intron 1 of a gene similar to hnRNP core protein A1 from Chromosome 7 (13510 EID identifier) have a homologous 250-nt internal region within much longer introns. That this is a low-level repeat is again reinforced by the finding of the same sequence on Chromosomes 1, 7, 8, 9, 11, and X. An online BLAST search of GenBank also revealed strong homology to a fragment of primate mtDNA.

Another class of homologous intron pairs is illustrated in Figure 3. In this instance, we found that the 145-nt long second intron of ABC-transporter, an evolutionarily conserved protein found on Chromosome 1, has 97% identity (256 BLAST score bits) to the first intron of a hypothetical gene on Chromosome 22. Amino acid sequences coded by the flanking exonic sequences show no homology. However, comparison of the nucleotide sequences of these genes revealed that nucleotide homology extends into the neighboring exonic and noncoding sequences (see Fig. 3). This is caused by a frameshift in the hypothetical gene. The revealed intron homology exists because of recent duplication of a region of at least 8 kb including the entire ABC-transporter gene into a new location on Chromosome 22, not to an intron transposition.

There are several such cases. We found homologous regions inside an intron of RNA-binding protein (6416 EID identifier) on Chromosome 3 and an intron in a hypothetical gene (31801 EID identifier) on Chromosome Y. Here, as well, the homologous area is not confined to intronic sequences but includes neighboring exons (again translated in different reading frames in the two genes). In 80 of the 118 investigated cases, one of the homologous introns was found in a hypothetical gene that encodes a protein with no significant similarity to any known peptide chains. That the existence of these genes as real coding units is dubious has been recognized in their removal (or drastic change in exon-intron structure) in more recently revised computer annotations of the human genome provided by NCBI.

In sum, in our entire analysis of the complete human genome, we found no example of homologous introns in non-homologous genes that showed evidence for an intron transposition event.

Comparison of *A. thaliana* Introns

We used 106,902 introns of *A. thaliana* for our analysis. Because plant introns are generally much shorter than those of human (all introns are <7 kb), and do not contain as many pseudogenes and DNA-repeats, it was feasible to compare all of them with each other without their prior division into subsets of similar-size introns. Also, because *Arabidopsis* has a much smaller genome than human, we were able to use the less stringent threshold for intron homology of a BLAST score of 65 bits. The main results of our analysis are presented in Table 1. We compared 1.1×10^{10} intron pairs. For 6971 pairs, the similarity score was above 65 bits (e -value $<10^{-10}$). The 6633 intron pairs that came from homo-

Table 1. Results of All BLAST Intron Comparisons

Species	Total number of analyzed intron pairs	Number of pairs with homology detected by BLAST	Number of pairs from homologous genes (discarded)	Number of pairs from same contig (discarded)	Final number of remaining intron pairs
Human	2.1×10^8	13,435	12,067	1250	118
<i>Arabidopsis thaliana</i>	1.1×10^{10}	6971	6633	169	169
<i>Drosophila melanogaster</i>	7.8×10^8	4073	3625	381	67

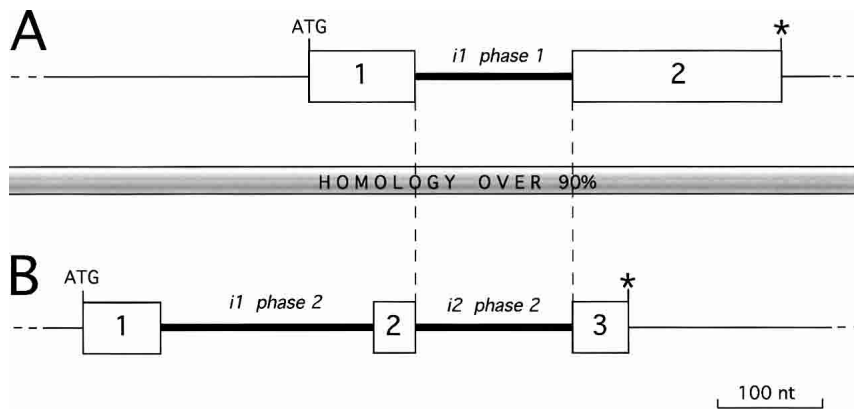


Figure 3 Comparison of (A) a hypothetical gene on Chromosome 22 (29720 EID identifier) at the top of the figure and (B) a ABC-transporter-like gene from Chromosome 1 (1786 EID identifier) shown at the bottom. The coding parts of the exons are shown as boxes, introns as thick lines, and noncoding regions as thin lines. Start and stop codons are shown as ATG and *, respectively. BLAST comparison revealed 97% identity between the second phase 2 intron of the ABC-transporter and the first phase 1 intron of the hypothetical protein. Manual examination of these genes showed that nucleotide identity (95% overall) extends over the entire shown region and beyond, extending at least 8000 nt. Despite the fact that the nucleotide sequences of exons 2 and 3 of the ABC-transporter are homologous to exons 1 and 2 of the hypothetical gene, they code nonsimilar polypeptides because they are translated in different frames shifted by 1 nt.

scription of that mRNA and subsequent recombination of the new intron-containing DNA with its intronless genomic original (Sharp 1985). Under this model, the rate of reverse transcription, which varies across species, should considerably influence the rate of intron acquisition. Yet, reverse transcription is very common in human. Thousands of processed pseudogenes are found throughout the human genome (Harrison and Gerstein 2002), and several cases of de novo insertions of retrotransposons have even been described (Ostertag and Kazazian 2001). Therefore, the absence of recent intron transpositions in the human genome is not easily reconcilable with intron movement via reverse splicing—reverse transcription—recombination.

Another proposed scenario postulates that spliceosomal introns originated from group II introns—a small set of very ancient self-splicing introns found in prokaryotes and in mitochondrial and chloroplast eukaryotic genomes (Roger and Doolittle 1993; Cavalier-Smith 1985; Martinez-Abarca and Toro 2000). The mechanisms of splicing and consensus sequences of group II introns are very similar to spliceosomal introns, indicating that these two types of introns probably have a common origin. Group II introns are mobile elements and transpose from one gene to another via ectopic retrotransposition that involves the reverse splicing of intronic RNA directly into DNA target sites (Dickson et al. 2001). The absence of group II introns in animals is congruent with the lack of intron transpositions during recent human and fruit fly evolution. However, specific spliceosomal proteins along with spliceosomal introns were recently found in all studied unicellular and early-branched eukaryotes, indicating that spliceosomal introns already existed at the time of eukaryotic origin (Fast et al. 1998; Fast and Doolittle 1999; Douglas et al. 2001; Archibald et al. 2002; Nixon et al. 2002). Hence, the hypothesis of recent evolutionary transformation of group II introns into spliceosomal ones seems rather simplistic and requires additional support.

There is one documented event of the de novo spliceosomal intron acquisition in the *Sh2* gene encoding one subunit of ADP-glucose pyrophosphorylase in maize (Giroux et al. 1994). This intron acquisition occurred via insertion of the DNA-transposable element *Dissociation (Ds)*, which mutated and became a new intron soon after the insertion. Because it is only one

example of such a mechanism of intron acquisition, it could be the exception rather than the rule in intron gain. Nonetheless, this mechanism of intron gain is attractive. There are dozens of different types of transposable elements in the genomes of animals and plants. These elements are not invariable—occasionally new types appear during evolution, whereas some old types, which have lost their ability to move, are degrading and vanish from genomes. A chance appearance of a new type of transposable element with GT and AG dinucleotide termini at its 5' and 3' ends, respectively (thus providing intron consensus splice sites), would create an invisible transposon. After the transposition of such elements into genes, the splicing machinery could splice them out of pre-mRNA transcripts, thus leaving the flanking coding sequence intact and creating a new intron at a previously intronless position, allowing for the rapid propagation of introns. Thus, during a short period of time, the intron gain of these intron-resembling transposons could be very intensive. Afterward, when this transposable element became inactive, the intron gain process could come to a complete halt. However, the sign of this event would be a large number of highly homologous introns of the same length, until this pattern was washed out by evolution. Clearly, there is no such event in the primate radiation.

Finally, it is important to note that we have only tested one model for the creation of new introns. Tests of the evidence for other models of intron origin, for instance, the model of Rogers (1989) and Venkatesh et al. (1999), in which tandem duplication of an exon or gene gives rise to novel introns through the use of cryptic splice sites, await further examination.

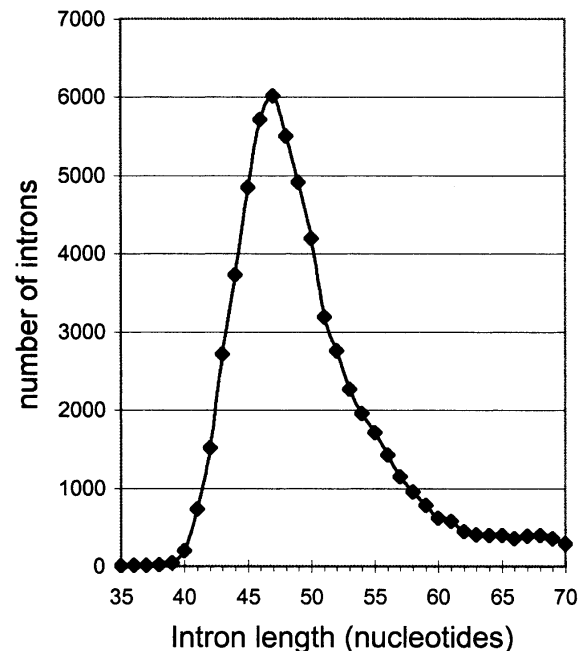


Figure 4 Distribution of *Caenorhabditis elegans* introns by length.

To understand the real mechanism of intron acquisition, we must find and analyze several examples of recently acquired introns. Such cases, which will involve the appearance of novel sequence within a phylogenetic pattern, would shed light on the question of intron gain.

METHODS

Analysis of Human Introns

The human Exon–Intron database (EID) was prepared using the EID program package (Saxonov et al. 2000) on the basis of the human chromosome sequences (“hs_chrN.gbk” files, $N = 1, 2, \dots, 22, X, Y$), downloaded from GenBank (Benson et al. 1999) on 12/20/2001. This initial Human EID consisted of 31,833 intron-containing genes. We checked all of the genes for the validity of the reading frame and removed 2264 genes that have internal stop codons. The rest of the 29,569 genes were used to generate flat fasta-formatted databases of exon sequences (188,881 entries), intron sequences (159,312 entries), and protein sequences (29,569 entries). Furthermore, we removed extra large introns and obtained 147,796 intron sequences with length $< 20,000$ nt. We found that 40% of the intronic sequences represent DNA repetitive elements using the RepeatMasker program of A.F.A. Smit and P. Green (<http://repeatmasker.genome.washington.edu>). All these repetitive sequences inside introns were automatically masked (replaced by Ns).

Because both 5' and 3' termini of introns are crucial for their splicing, it is reasonable to assume that a newly inserted intron should have the same length as its original copy. This assumption allows us to considerably simplify the computer analysis of intron homology by examining only those intron pairs with comparable lengths. Intronic sequences were divided into two sets of batches of introns of similar size. Set one consists of the following: batch 0 (introns with lengths ≤ 50 nt), batch 1 (51–100 nt), batch 2 (101–150 nt), and so on. Set two has a shift of 25 nt, containing batches of introns with lengths of 26–75 nt, 76–125 nt, 126–175 nt, and so on.

We used stand-alone gapped BLAST 2.0 (Altschul et al. 1997) to do pairwise comparisons for each pair of introns within each batch. Those intron pairs that gave BLAST-scores over 100 bits were selected for further consideration (for introns < 150 nt, we used a lower homology threshold of 80 bits). We chose this threshold as sufficiently stringent to ensure common ancestry but sufficiently lenient to detect homologies in pairs of short sequences diverged millions of years ago. For example, the first introns of the β -globin genes of human and brown lemur (the species diverged ~ 25 Mya) are 130 nt long and have 86 identical bases in a 103-nt-long region. The BLAST score for this alignment is exactly 100 bits (probability e -value is 4×10^{-19}). The selected intron matches were narrowed down by eliminating those that belong to genes coding for homologous proteins (see below). In addition, we removed those pairs of homologous introns in genes found very close to each other on the same chromosomal locus (belonging to the same contig, that is, deriving from the same GenBank file). Pairs of introns that satisfied this series of criteria were then inspected manually.

Protein Groups

All 29,569 human proteins were compared with each other by stand-alone BLAST 2.0 binaries (Altschul et al. 1997). Next, we performed the simplest grouping procedure: (1) two proteins were considered homologous and put in the same group if they have homology above 55 bits; (2) groups were pooled together if any member of one was homologous (again, at 55 bits) to any member of the other. This procedure yielded 17,054 different protein groups.

Analysis of Animal and Plant Introns

D. melanogaster, *C. elegans*, and *A. thaliana* genome sequences were downloaded from the most recent version of GenBank. Exons and introns were characterized by the EID program package

(Saxonov et al. 2000). As a result, we obtained 106,902 *Arabidopsis* introns, 39,624 *Drosophila* introns, and 122,490 *C. elegans* introns. *Drosophila* and *Arabidopsis* introns were analyzed in the same manner as human introns, described above. The only difference consisted of not dividing *Drosophila* and *Arabidopsis* introns into batches of similar sizes. Our analysis of *C. elegans* introns was slightly different. There is a very large number of *C. elegans* introns with a length of 47 nt, constituting a very sharp and very narrow peak in the *C. elegans* intron length distribution. All 6021 47-nt-long introns were collected and analyzed according to their similarity with each other by a special program designed for this task.

All calculations were performed with computer programs written in Perl on a LINUX platform workstation with dual Pentium processor. The described databases are available on our Web site: <http://www.mcb.harvard.edu/gilbert/eid>.

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