COMMUNICATION

Alternative splicing and genomic stability

Kevin Cahill

Department of Physics and Astronomy, University of New Mexico, Albuquerque, NM 87131, USA and
Center for Molecular Modeling, Center for Information Technology, National Institutes of Health,
Bethesda, MD 20892-5624, USA
E-mail: cahill@unm.edu

Received 7 March 2004
Accepted for publication 10 June 2004
Published 30 June 2004
Online at stacks.iop.org/PhysBio/1/C1
doi:10.1088/1478-3967/1/2/C01

Abstract
Alternative splicing allows an organism to make different proteins in different cells at different times, all from the same gene. In a cell that uses alternative splicing, the total length of all the exons is much shorter than in a cell that encodes the same set of proteins without alternative splicing. This economical use of exons makes genes more stable during reproduction and development because a genome with a shorter exon length is more resistant to harmful mutations. Genomic stability may be the reason why higher vertebrates splice alternatively.

For a broad class of alternatively spliced genes, a formula is given for the increase in their stability.

What is alternative splicing? A procaryote (no nucleus) transcribes one or more genes into mRNA and immediately translates the mRNA into protein. But a eucaryote first transcribes a single gene into pre-mRNA, and then, using spliceosomes, turns the pre-mRNA into mRNA by splicing out most or all of its introns and often many of its exons. The eucaryote then exports the mRNA out of its nucleus into its cytosol, where its ribosomes translate the mRNA into protein. A eucaryote can often make different proteins from the same pre-mRNA transcript by splicing it in different ways. This trick is called alternative splicing.

Why do higher vertebrates splice alternatively? Alternative splicing allows an organism to make different proteins in different cells at different times, all from the same gene, by poorly understood regulatory devices (Alberts et al 2002). But this diversity of proteins could also be produced by several different genes controlled by promoters and enhancers—in fact, that is how biologists thought genes worked until they discovered alternative splicing. The advantage of alternative splicing is that its economical use of exons makes genes more stable during reproduction and development.

This communication gives a formula and a rule of thumb for the increase in the stability of a broad class of alternatively spliced genes. The DSCAM gene of the fruit fly and the cSlo gene of the chicken provide examples that illustrate the formula and the rule. A Monte Carlo simulation, displayed in figure 1, suggests how alternative splicing may help dividing human cells avoid excessive mutations.

How does alternative splicing make genes more stable? Consider, for instance, a gene that has a long exon of 1000 base pairs (b) and two short ones, each 100 b long. The total length of its exons is 1200 b. Alternative splicing allows the cell to make two different RNAs, each of 1100 b long. Without alternative splicing, the cell would need two genes, each 1100 b long, for a total exon length of 2200 b. Thus in this example, alternative splicing reduces the length of the exons in the DNA by 45%. This reduction in the length of exonic DNA implies a reduction of 45% in the error rate during the replication of this gene. In effect, the gene is nearly twice as stable due to alternative splicing. Since an error in the replication of critical exonic DNA is potentially lethal, this extra genomic stability is biologically significant and is one of the reasons why higher eucaryotes use alternative splicing. Computer scientists will recognize alternative splicing as akin to file compression (Ford 2001).

More generally, let us consider a gene that has $M$ groups of mutually exclusive exons in addition to the constitutively
spliced exons (the exons that are always kept in the mRNA). For each group \( i (i = 1, \ldots, M) \), let \( N_i \) denote the number of mutually exclusive exons in the group, including the null exon of length zero if the spliceosome may splice out all the exons of the group. Assume that the spliceosome always selects at most one exon and no introns from each of the \( M \) groups of \( N_i \) exons with no shuffling. Assume that the organism expresses all possible proteins at some time in some cell.

Let us use \( L_c \) for the total length in nucleotides of the constitutively spliced exons. Without alternative splicing, these \( L_c \) nucleotides would be repeated in each of the \( N \) proteins for a total length of \( N L_c \).

If \( L_{ik} \) is the length of exon \( k \) of group \( i \), then without alternative splicing, each of the \( N_i \) exons of length \( L_{ik} \) would be repeated \( \frac{N}{N_i} \) times. So the total length devoted to group \( i \) without alternative splicing would be

\[
N = \prod_{j=1}^{M} N_j
\]

Thus, the number of nucleotides that would be needed to encode all \( N \) proteins and that would have to be copied correctly each time a cell divides is

\[
N_{nas} = N \left( L_c + \sum_{i=1}^{M} \frac{1}{N_i} \sum_{k=1}^{N_i} L_{ik} \right)
\]

without alternative splicing.

But with alternative splicing, the number of needed nucleotides is only the length of all the exons,

\[
N_{as} = L_c + \sum_{i=1}^{M} \sum_{k=1}^{N_i} L_{ik}
\]

Since the error rate in the replication of DNA is \( 10^{-9} \) per base pair (Alberts et al 2002), the probability of an exonic error in the gene during replication is \( N_{nas} \times 10^{-9} \) without alternative splicing, but only \( N_{as} \times 10^{-9} \) with alternative splicing. So if we ignore the critical control sequences in the introns, then the ratio

\[
I = \frac{N_{nas}}{N_{as}}
\]

is the increase in the stability of the gene due to alternative splicing. The intron control sequences probably boost \( I \) slightly.

The DSCAM gene of Drosophila provides a striking example of alternative splicing. This gene encodes receptors that guide the growth of the axon of Bolwig’s nerve in the fly embryo (Schmucker et al 2000). It has \( M = 4 \) groups of \( N_1 = 12, N_2 = 48, N_3 = 33, \) and \( N_4 = 2 \) exons (Schmucker et al 2000, Black 2000). The exons in each group are mutually exclusive, and the total number of possible proteins is

\[
N = 12 \times 48 \times 33 \times 2 = 38016.
\]

The DSCAM gene, including introns, is 61.2 kb long, and its mRNA, after transcription and splicing, contains 24 exons and is 7.8 kb long (Schmucker et al 2000, Black 2000).

The ratio \( N_{nas} / N_{as} \) depends explicitly upon the lengths \( L_c \) and \( L_{ik} \). Since most internal exons are between 50 and 300 nucleotides in length (Smith and Valcárcel 2000), let us simplify the bookkeeping by setting \( L_{ik} = 200 \) b. The spliced DSCAM mRNA is 7.8 kb long and contains 4 alternatively spliced exons and 20 constitutively spliced exons. So the set of constitutively spliced exons is of length

\[
L_c = 7800 - 4 \times 200 = 7000
\]

or \( L_c = 7 \) kb. Thus by equation (4), the exonic length required with alternative splicing is

\[
N_{as} = L_c + 200 \sum_{i} N_i
\]

\[
= 7000 + 200(12 + 48 + 33 + 2) = 26000
\]
or \( N_{\text{as}} = 26 \text{ kb} \). But by equation (3), the exonic length required without alternative splicing is

\[
N_{\text{as}} = N(L_c + 200 M)
\]

\[
= 38,016 \times 7800 = 296,524,800
\]

(8)

or \( N_{\text{as}} = 297 \text{ Mb} \), which, incidentally, is nearly twice the length of the entire *Drosophila* genome and about six times the length of all the exons in the human genome. With these assumptions, the chance of a crucial error in the DSCAM gene during replication is 0.30 without alternative splicing, but only 2.6 \( \times \) \( 10^{-5} \) with alternative splicing. The ratio

\[
I = \frac{N_{\text{as}}}{N_{\text{as}}} = 11,400 = 1.1 \times 10^4
\]

(9)

is the increase in genetic stability due to alternative splicing.

Fruit flies without alternative splicing would accumulate about 10,000 exonic DSCAM errors in 30,000 generations (2500 years), and each fly would have its own set of 10,000 errors. Over this period, the DSCAM gene of the fly population gradually would become uniformly dysfunctional with relatively small differences in fitness among individual flies. On the other hand, flies with alternative splicing would accumulate less than one exonic DSCAM error in 30,000 generations. Moreover, the probability that the one error would occur in the \( L_c \) exons that are constitutively expressed would be \( L_c/N_{\text{as}} = 7/260 = 0.27 \), and that unlucky fly would be distinctly unfit. Thus, alternative splicing not only avoids exonic errors; it also helps natural selection weed out unfit individuals. Alternative splicing and natural selection cooperate to preserve the integrity of the genome.

In most genes, the increase in genomic stability due to alternative splicing might be like 5 or 10 than \( 10^5 \), but even a 500% increase in genetic stability during reproduction and development is worth the trouble of alternative splicing. If without alternative splicing the average genes were five times longer, then 7.5% rather than 1.5% of the genomes of flies. The DNA of a human diploid cell has 6.4 billion base pairs. The error rate of \( 10^{-9} \) per base pair implies that on average there will be 6.4 errors per cell division. With alternative splicing, only 1.5% of these errors occur in exons and are potentially deleterious, so the probability of a daughter cell with perfect exons is approximately \( P_{\text{as}} = 1 - 6.4 \times 0.015 = 0.904 \). A more accurate estimate is

\[
P_{\text{as}} = (1 - 10^{-9})^{0.015 \times 6.4 \times 10^9} \approx e^{-0.096} = 0.908
\]

(10)

Without alternative splicing, 7.5% of the errors would occur in exons, and so the probability of a daughter cell with perfect exons would be roughly \( P_{\text{as}} = 1 - 6.4 \times 0.075 = 0.52 \). A more accurate estimate is

\[
P_{\text{as}} = (1 - 10^{-9})^{0.075 \times 6.4 \times 10^9} \approx e^{-0.48} = 0.619
\]

(11)

The adult human arises from about 46 cell divisions, so the probability that any given adult cell has perfect exons is \( (P_{\text{as}})^{46} = 0.012 \) with alternative splicing, but only \( (P_{\text{as}})^{46} = 2.6 \times 10^{-10} \) without alternative splicing.

To estimate the implications of alternative splicing for human evolution and development, I again assumed that the human genome without alternative splicing would have five times more exonic base pairs. I let two sets of 1000 cells divide 50 times *in silico*. The set of cells that used alternative splicing had 0.015 \( \times \) 6.4 \( \times \) \( 10^9 \) = 9.6 \( \times \) \( 10^7 \) exonic base pairs; the set that did not use alternative splicing had five times as many or \( 4.8 \times 10^8 \) exonic base pairs. I divided the 1000 cells into 20 groups of 50 cells each and plotted in the figure the average number of exonic errors per cell for each of the 20 groups with and without alternative splicing. As shown in figure 1, the average number of defective exonic base pairs per daughter cell after 46 cell divisions is 4.43 \( \pm \) 0.07 with alternative splicing (lower, green lines) but 22.0 \( \pm \) 0.14 without alternative splicing (upper, red lines). Since with alternative splicing, cells free of exonic error produce daughter cells that also are free of exonic error at a rate of 91%, apoptosis followed by division of adjacent cells can correct the 1 or 2 of the 4 exonic errors that are troublesome. But because without alternative splicing, cells free of exonic error produce daughter cells free of exonic error at a rate of only 62%, it is hard to see how apoptosis could cope with 22 exonic errors per adult cell.

We may derive a rule of thumb for the increase in genetic stability by noting that \( L_c \) defined by

\[
\langle L_c \rangle = \sum_{i=1}^{M} \frac{1}{N_i} \sum_{k=1}^{N_i} L_{ik}
\]

(12)

is an effective average length of the alternative exons that are spliced into the mRNA and that \( \langle N \rangle \) defined by

\[
\langle N \rangle \langle L_c \rangle = \sum_{i=1}^{M} \frac{N_i}{L_{ik}}
\]

(13)

is a kind of average of the numbers \( N_i \) of alternative exons in the \( M \) groups. Let us further use \( r \) for the ratio of the average length \( \langle L_c \rangle \) of the selected exons to the length \( L_c \) of the constitutively spliced exons

\[
r = \frac{\langle L_c \rangle}{L_c}
\]

(14)

Then with these definitions, the increase \( I \) in genetic stability is

\[
I = \frac{N_{\text{as}}}{N_{\text{as}}} = N \left[ 1 + r + (N/r) \right]
\]

(15)

The fraction that multiplies the total number \( N \) of possible proteins is less than unity. But it is generally not tiny because the ratio \( r \) usually is small and because \( N \) usually is less than 30. In the case of *Drosophila* DSCAM and with the assumptions \( L_c = 7.0 \text{ kb} \) and \( L_{ik} = 200 \text{ b} \), the four selected exons are of length \( \langle L_c \rangle = 800 \text{ b} \); the ratio \( r = 800/7000 = 0.114 \); and the effective average number \( \langle N \rangle \) of exons per group is \( (N) = 95 \times 200/\langle L_c \rangle = 95/4 = 23.7 \). The fraction \( (1 + r)/(1 + (N/r)) = 3/10 \), and the increase in genetic stability is \( I = 0.3 N = 11,400 \).

Hearing in chickens provides another example of the contribution of alternative splicing to genetic stability. The \( cSlo \) gene of the chicken cochlea encodes the membrane proteins that form the Ca\(^{2+}\)-activated K\(^+\) channels that determine the resonant frequency of each hair cell in the basilar papilla. Alternative splicing provides some \( N = 576 \) variants of the mRNA for each of the four components of this tetramer.
membrane protein (Rosenblatt et al 1997, Navaratnam et al 1997, Black 1998), resulting in a huge number of possible resonant frequencies. In cSlo, the ratio $r = 0.1$, and the mean number $\langle N \rangle$ of exons in each of the eight groups is about 2.6 (Rosenblatt et al 1997). So by the rule of thumb (15), alternative splicing increases the genetic stability of cSlo by a factor of about

$$I = \frac{576^{1.1}}{1.26} = 503.$$  \hspace{1cm} (16)

The tetrameric structure of the functional membrane protein effectively boosts $I$ by another factor.

Another example of alternative splicing’s exonic economy is provided by the mammalian immune system, which uses site-specific genetic recombination in developing B cells (Alberts et al 2002).

We have seen that the exonic economy of alternative splicing increases the stability of the genome. As genomics and proteomics advance, the protein-to-gene ratios of the higher vertebrates will teach us how much alternative splicing actually contributes to the stability of their genomes.

Acknowledgments

Thanks to Mark Fleharty, Michael Malik, Valerie Parsegian, Stephanie Ruby, and James Thomas for helpful conversations. Peter J Steinbach kindly hosted the author during his sabbatical at NIH.

References

Alberts B, Johnson A, Lewis J, Raff M, Roberts K and Walter P 2002 Molecular Biology of the Cell 4th edn (New York, NY: Garland Science)
Black D L 1998 Neuron 20 165–8
Black D L 2000 Cell 103 367–70
Ford J L 2001 The Book of Winzip [Electronic Resource]: File Compression and Archive Management Made Easy (San Francisco, CA: No Starch Press)
Navaratnam D S, Bell T J, Tu T D, Cohen E L and Oberholtzer J C 1997 Neuron 19 1077–85
Rosenblatt K P, Sun Z-P, Heller S and Hudspeth A J 1997 Neuron 19 1061–75
Schmucker D, Clemens J C, Shu H, Worby C A, Xiao J, Muda M, Dixon J E and Zipursky S L 2000 Cell 101 671–84
Smith C W J and Valcárcel J 2000 Trends Biochem. Sci. 25 381–8