What controls the length of noncoding DNA?
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Several recent studies of genome evolution indicate that the rate of DNA loss exceeds that of DNA gain, leading to an underlying mutational pressure towards collapsing the length of noncoding DNA. That such a collapse is not observed suggests opposing mechanisms favoring longer noncoding regions. The presence of transposable elements alone also does not explain observed features of noncoding DNA. At present, a multidisciplinary approach — using population genetics techniques, large-scale genomic analyses, and in silico evolution — is beginning to provide new and valuable insights into the forces that shape the length of noncoding DNA and, ultimately, genome size. Recombination, in a broad sense, might be the missing key parameter for understanding the observed variation in length of noncoding DNA in eukaryotes.

Introduction

The total amount of DNA in a haploid genome, known as the C-value, varies enormously among eukaryotic organisms, ranging from 1.2 × 10^7 bp (e.g. the yeast Saccharomyces cerevisiae) to >6 × 10^11 bp (e.g. Anoetha dioica). More intriguingly, this irregularity is also frequently observed between closely related species (i.e. within the same taxonomic group or genus) with similar levels of complexity (morphological, developmental, behavioral, etc.), number of genes, and regulatory networks. As emblematic examples, the size of genomes of flowering plants varies 1000-fold, and the DNA content can change >200- and 100-fold among vertebrates and species of salamanders, respectively [1,2]. The human genome size is not exceptional among vertebrates, with a genome larger than that of birds but smaller than those of most fishes or amphibians; mammalian genomes vary by >40% in length [3]. Drosophila species provide another interesting case, where differences in genome size show no obvious relationship to rate of cell division or development time, nor do they show clear phylogenetic trends [4], indicating that the forces involved in genome size change may act quite rapidly. This lack of correspondence between genome size and biological complexity has been called the C-value paradox or enigma [1,5,6].

Eukaryotic genomes consist mostly of DNA sequences that are not part of coding regions, regulatory elements, or RNA genes, and it is this apparently superfluous DNA that contributes to the variation in C-value. Most euchromatic noncoding DNA comprises introns and intergenic regions, with repetitive sequences of different types — satellite and microsatellite DNA, and different classes of transposable elements (TEs). In this review, I begin by describing several genome features of euchromatic noncoding DNA, including intron and intergenic length, and TE and microsatellite presence, focusing particularly on their relationship with recombination rates across genomes. Next, I detail results of studies on the mutational tendencies of small indels (insertions or deletions). Finally, I discuss selective forces that may be influencing the length of noncoding regions.

Genome features

Intron size

In an innovative study, Lengel and Penman [7] showed that the size of hnRNA, but not mature mRNA, increases with genome size in dipterans. They reported significantly longer hnRNA in Aedes than in Drosophila, consistent with Aedes’ 5–6-fold larger genome. This observation, dated before the discovery of the intervening sequences or introns in 1977, was the first indication of a positive relationship between genome size and total intron length. A significant, although weak, positive relationship between intron and genome size has now been established for many other eukaryotes [8–11], both on a large evolutionary timescale [11] and between Drosophila species [9]. In all cases, however, the differences in intron size alone cannot fully account for the differences in euchromatic genome size, indicating that the differences in genome size are not easily explained by a single class of noncoding DNA. Variation in genome size among organisms is usually associated to congruent changes across different classes of noncoding DNA (e.g. introns and intergenic regions), suggesting that they may be responding to similar, or at least overlapping, evolutionary forces. Two factors that can influence the length of introns and intergenic regions are TE and, to a lesser degree, microsatellite presence.

Transposable elements

TEs are ubiquitous in all eukaryotes and can be a leading factor influencing the length of noncoding DNA and genome size in some species. For instance, in humans, recognizable TEs represent a fraction as large as 45% of the euchromatic DNA, and are abundantly present in introns [12,13]. In other organisms, such as Drosophila melanogaster, TEs have a more limited influence on the overall size of noncoding sequences, with minor effects on intron length; only 0.4% of introns in genes with known full-length mRNAs have detectable remnants of TE sequences (JM Comeron, unpublished data). The rare detection of TEs in Drosophila...
intron length is not unexpected because they may often alter the accuracy of transcript processing, mostly with deleterious consequences on fitness. In organisms with much longer introns (e.g. humans), the insertion of TEs may have an attenuated effect on gene expression and would further increase the chance of successive insertions. An equivalent argument can be made for intergenic regions based on the relationship on gene expression of long terminal repeats in the proximity of a gene.

TE invasion and expansion can cause rapid changes in genome size, generating differences between very closely related species that might share most other forces that influence genome size. An extreme example is the genome of maize, which has doubled its size as a result of retrotransposon insertions, mostly in the last three million years [14]. In *Drosophila*, the TE distribution can also vary among closely related species, or even among populations of the same species [19•–21•]. The recent invasion by the Penelope TE of *D. viridis* [16•] is an interesting case in point.

**Microsatellites**

The main mutational mechanism causing changes in the number of microsatellite repeats is polymerase template slippage that is not corrected by the mismatch repair system [17,18]. In *Drosophila*, recent studies have revealed differences both in density and length of microsatellites among species [19•–21•]. These differences concur with the idea that larger genomes also tend to have more and longer microsatellites [22]. Among distant organisms, *Saccharomyces cerevisiae*, *Caenorhabditis elegans*, *Arabidopsis thaliana*, *D. melanogaster*, and *Homo sapiens* show a congruent increase in overall microsatellite presence with genome size when both length and density of microsatellites are considered [23–25,26•,27]. This relationship, however, is not universal, with the pufferfish *Fugu rubripes* having a higher density and longer microsatellites than humans but with a genome eight times smaller [28]. Nevertheless, the general trend suggests that the forces acting on intron and intergenic length are not independent of those influencing abundance and length of microsatellites.

**Noncoding regions and recombination rates across genomes**

The study of characteristics of noncoding DNA in relation to recombination rates across a given genome is informative because population genetics models of selection predict that recombination increases the efficacy of selection [29,30]. Therefore, assuming that there is no bias in the indel-repair mechanism associated with recombination (see below), the observation of different patterns of noncoding DNA between regions of high and low recombination may reveal the varying action of selection.

**Introns and intergenic regions**

In humans, there is evidence [13••,31,32••] that introns are shorter in isochores with high recombination (i.e. G+C-rich isochores [33,34]) than in those isochores with low recombination rates (i.e. G+C-poor isochores). Moreover, gene density and the length of intergenic regions also decrease with recombination [13••], suggesting qualitatively similar forces shaping the length of both types of noncoding DNA across genomes.

Further, the observed relationship between the length of noncoding DNA and recombination in the human genome can be only partially attributed to TE presence. For instance, both median intron length and gene density vary ~10-fold between GC-rich and GC-poor genomic regions but the proportion of the genome comprising TEs varies much less conspicuously between these regions [13••,35]. It is possible to argue that the limited variation of TE presence is caused by the fact that TEs in regions with low recombination are older than in regions with high recombination; they therefore tend to be no longer recognizable but still contribute to size. This possibility is unlikely in humans, however, where the opposite trend is observed, with TEs younger in GC-poor isochores [13••], ruling out a simple mutational scenario with rare TE insertions and frequent small deletions [36,37] to explain variation in length of noncoding DNA in humans.

In *Drosophila*, introns also tend to be longer in regions of low recombination [32••,38••]. This relationship is mostly caused by genes with long coding sequence (JM Comeron, unpublished data) and is not explained by TE presence alone (see above). Congruently, gene density also increases with recombination in *D. melanogaster* (JM Comeron, unpublished data), paralleling the relationship between the length of noncoding regions and recombination observed in the human genome. A rare case of very long introns can be found also in *Drosophila*, where genes located in the heterochromatic Y chromosome can have mega-introns (>1 Mb) with large clusters of satellite DNA [39,40•]. This result suggests either that recombination is required to prevent a one-way growth process for satellite DNA or that selection against very long introns is inefficient in this genomic region.

**Transposable elements**

Although TEs are not the only contributors to differences in length of noncoding DNA across a given genome, they may play some role. There are at least two reasons to suggest that TEs should accumulate in chromosomal regions with reduced recombination. First, non-homologous recombination between different elements has deleterious consequences on fitness because it induces chromosomal rearrangements [41]. Second, as indicated above, selection against the deleterious effects of TEs either in or near genes is more efficient in regions with high recombination. In *Drosophila*, this general pattern is observed when the frequency of a given element in the population is taken into account [41,42]. In addition, different TE elements might show characteristic distributions across genomes [15•], representing different stages in the temporal dynamics of TE invasion. Illustrative of this pattern is the
Penelope element. In *D. virilis*, the recently inserted Penelope sequences are restricted to recombining euchromatic regions whereas in other species of the virilis group, where the element has been long present, Penelope is mainly detected in heterochromatic regions [43].

Interestingly, recent studies have shown that recombination rates in particular genomic regions of *Drosophila* can change rapidly. For instance, the X-telomeric region in *D. melanogaster* has strongly reduced frequency of crossing over compared to other genomic regions. The comparison with other species of the melanogaster group shows that this reduced crossover frequency was recently acquired in the *D. melanogaster* lineage, after the evolutionary split between *D. melanogaster* and *D. yakuba* [44*]. Incidentally, this finding might suggest that TEs may not have yet reached the new equilibrium of presence and segregating frequency in this genomic region in *D. melanogaster*, which might explain the absence of high TE accumulation at the tip of the X chromosome [41,42].

*Caenorhabditis elegans* is an intriguing case also worth noting. *C. elegans* shows the pattern opposite to that in *Drosophila* or humans, with a positive relationship between TE density and recombination rate [45,46], although some miniature inverted-repeat TE (MITEs) show different distribution profiles [47]. This pattern might suggest that selection against TEs is not an important factor explaining TE distribution in *C. elegans* [46]. It might also suggest that regions with high recombination in *C. elegans* are, overall, under reduced selective constraints, congruent with a low density of both highly expressed genes and conserved eukaryotic genes [45]. Alternatively, it could indicate a recent TE invasion, mostly targeting regions of high recombination, hence causing unusual and unstable genomic features.

### Microsatellites

Genome analyses indicate that the differences in recombination rate across *D. melanogaster*'s genome has no effect on either microsatellite density or average repeat number [25]. This result is a first indication that there is no directional bias associated with recombination-dependent indel mismatch repair (see below).

### Mutational tendencies of small indels

Two different approaches have been used to study the mutational rates and tendencies of indels: first, the analysis of indels in pseudogenes and DOA elements, and second, the study of polymorphic indels in noncoding regions.

### Indels in pseudogenes and DOA elements

The study of pseudogenes and DOA elements is based on the assumption that these sequences are free of selective constraints for coding information and, therefore, the observed indel patterns are a faithful representation of the indel mutation process. Early studies, mostly of mammalian pseudogenes, showed that small deletions are more frequent than insertions [48-50]. The study of 603 small indels in 156 processed pseudogenes from humans and murids shows a ratio of deletions to insertions (i.e. deletion bias [DB]) of 2.74 [50], evidence of a significant pressure towards DNA loss. Although DB is slightly larger in humans than in murids (2.9 versus 2.6, respectively), counter to expectations based on the smaller size of the mouse and rat genomes compared to human [3], insertions are longer and deletions are shorter in the human lineage, and the overall rate of DNA loss (overall deletion bias [ODB]) is lower in the human lineage (1.69) than in mouse and rat (2.63).

A novel contribution to understanding the role of indel evolution on genome size involved studies of indels in nontransposing copies of non-LTR retrotransposable elements (i.e. DOA elements). These studies first focused on the Helena element in *Drosophila* [51,52] and on the Lau1 element in Hawaiian crickets of the genus *Laupala*, the latter having a genome 11 times larger than *Drosophila* [53*]. The rate of DNA loss in DOA elements differs between these insects, with a much higher rate in *Drosophila* (DB = 8.7 [52]) compared to *Laupala* (DB = 2.7 [53*]), in agreement with *Drosophila*'s more compact genome. Also in insects, the study of indels in 58 paralogous pseudogenes in *Podisma pedestris* [54*•*], which has a genome >100 times as large as that of *Drosophila*, shows a DB of 2.7 when hot spots for indel mutations are excluded from the analysis. ODB is also strongly affected by the size of indels in these species, with ODB of 74.7, 3.8 and 3.6, for *Drosophila, Laupala* and *Podisma*, respectively.

The study of indels in a large pseudogene family in *C. elegans* [55] also reveals an excess of deletions compared to insertions (DB = 3.8), with the detection of very large insertions and deletions. Intriguingly, the size of indels would generate an unexpected overall gain of DNA (ODB = 0.49 caused by two very long insertions) but no definitive conclusions can be made because the length of some long deletions may be gross underestimates [55].

The conclusion of these studies based on indels in pseudogenes and between DOA elements is that, overall, the rate of DNA loss is greater than that of DNA gain, at least based on small indels. Moreover, this rate of overall DNA loss is greatest in *Drosophila* compared to *Laupala, Podisma*, and mammals, congruent with the smallest size of *Drosophila* genome compared to the other genomes. *C. elegans* depicts a contrasting observation, with a very compact genome that is smaller than that of *D. melanogaster* but with a lower rate of DNA loss.

### Polymorphic indels

Weak selection has considerably less influence on the presence of mutations as polymorphisms in a population than it does on their probability of fixation [56] (Figure 1). Therefore, studies based on polymorphism presence may give us a pattern very close to the mutational tendencies,
Moreover, this study in D. melanogaster [32••] found equivalent pDBs in genomic regions with high and low recombination, suggesting that the observation of different lengths of noncoding regions in genomic regions with different recombination rates is not caused by a DB that changes with recombination rate. Conversely, the discrepancy between genomic patterns and indel polymorphisms unveils a possible role of selection in shaping the length of noncoding regions, at least in D. melanogaster. Future studies differentiating euchromatic versus heterochromatic, high versus low recombination regions, and fixed versus polymorphic indels, in DOA elements, pseudogenes and other noncoding regions will be useful for completing the picture of the rate of spontaneous DNA loss across the Drosophila genome.

**Directionality in indel repair mechanism**

Subtle differences in both the efficacy and the directionality of the indel mismatch repair system can generate significant differences in DB and in the length of noncoding DNA between species. Moreover, directionality in the recombination-dependent indel mismatch repair can generate patterns of noncoding length correlated with recombination rates across genomes. In yeast, however, there is no clear directionality in indel repair. The repair of mispaired loops in heteroduplex DNA shows diverse biases, efficiencies, and repair pathways, for short and long indels, for different genomic regions, and for nicked and continuous DNA (see references in [32••]). On the other hand, a generally biased repair towards deletions associated with recombination is not observed in D. melanogaster’s genome on the basis of available data of indel polymorphisms and their frequency in natural populations [32••], and microsatellite distribution [25]. Nonetheless, more definitive studies are required to appraise its actual contribution.

**Natural selection on the length of noncoding regions**

Irrespective of the precise rate towards DNA loss, this tendency will lead to the collapse of noncoding DNA, and TE insertion alone cannot explain the differences in length of noncoding regions between species or across genomes. Therefore, mutational explanations based on ‘junk’ [59] or ‘selfish’ [60] DNA don’t provide a sufficient explanation. Several selective hypotheses have been proposed to explain observed relationships between genome size and cellular and developmental traits — some observed only in particular lineages — although one cannot immediately discern between causal, indirect, and coincidental relationships. Typical of this class of selective hypotheses, the nucleotypic [61] and nucleoskeletal [62] theories propose structural functions to noncoding DNA, setting the minimum size attainable by a nucleus, or an indirect (or coevolutionary) response to genetically-controlled nuclear size.

Other selective explanations for the presence of noncoding DNA relate to its role in regulating gene expression [63].
Greater developmental and/or morphological complexity may require more elaborate regulatory systems, which in turn require larger amounts of noncoding DNA. Consistent with this idea, the comparison between the number of selectively constrained nucleotides per intergenic interval is distinctly higher in mammals than in C. elegans [64]. The increment in noncoding constrained sites, however, can explain only a small fraction of the difference in the length of noncoding regions between these organisms. This observation is another indication that the length of noncoding DNA is not solely determined by functional requirements.

Noncoding DNA as an enhancer of recombination

Theoretical and simulation studies [65–67] have shown that mutations that raise the recombination rate between strongly selected loci have an increased chance of fixation because they reduce the detrimental consequences of linkage between selected mutations. Results of in silico evolution (i.e. forward simulations) [68,69*,70*,71] have shown that many weakly selected mutations interfere with each others’ fixation probability, also reducing the efficacy of selection relative to expectations for independently evolving mutations (also known as the Hill–Robertson effect [72,73] or interference selection [IS]). IS increases with the number of sites under weak selection and is reduced with recombination.

Weakly selected mutations are numerous in natural populations and are physically clustered across genomes — mostly in genes, exons, and in regulatory regions — creating genomic regions with a high density of sites under selection, with limited recombination between them. This physical distribution of mutations causes IS to have a measurable effect in most eukaryotic genomes, and this raises the possibility that indels between clusters of selected sites (genes or exons) might be subject to natural selection as modifiers of recombination [32**]. Mutations that increase the distance (hence recombination) between mutations under selection will be favored by selection under this scenario because, ultimately, they increase the effectiveness of selection acting on these flanking sites. Longer introns and lower gene density will be favored in regions of low recombination. This tendency for selection to favor insertions and to oppose deletions will counter-balance the mutational bias towards DNA loss.

Genomic patterns in D. melanogaster and humans cannot be explained by either TE presence or small indels alone (see above). The fact that both intron length and intergenic sizes negatively correlate with recombination is consistent with the action of IS. In silico evolution results support detectable effects of small indels on the efficacy of selection in adjacent regions (JM Comeron, M Kreitman, unpublished data). Therefore, IS in recombining genomes might be influencing genomic features [32**] such as gene structure, intron length and gene density, and it might be key to the C-value paradox. Under this scenario, the length of noncoding regions represent a highly dynamic mutation–selection–drift equilibrium responding to changes in effective population size and/or recombination rates (per physical unit) as well as...
to variation of species-specific developmental requirements and mutational tendencies (Figure 2).

Conclusions

Indels are beginning to be studied using both population genetics and genomic techniques. As is the case for coding sequence evolution, these techniques and corresponding statistical tests should allow discrimination of the relative importance of selection, mutational tendencies, and random genetic drift in influencing the length of noncoding regions and genome size. This genomics-meets-population genetics approach can be implemented with the study of in silico evolution to capture the expected outcome of complex evolutionary interactions, including weak selection and drift. TE invasion and expansion, and a general mutational tendency towards DNA loss are two mutational forces that inescapably influence the length of noncoding regions in most eukaryotes. These mutational tendencies alone, however, cannot explain genomic features or indel polymorphism data, indicating that natural selection might be acting on indels to influence the length of noncoding regions. A recent proposal [32••] suggests that noncoding regions (introns and intergenic regions) can be viewed as modifiers of recombination, and thus have a selective role. Based on this suggestion, a better understanding of the C-value paradox might be attained when recombination is entered into the consideration, both as the population recombination parameter per site (ironically, often also noted by $C$ [74]), and as the physical distance between sites under selection.

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References and recommended reading

Papers of particular interest, published within the annual period of review, have been highlighted as:

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