

## NEWS AND COMMENTARY

Mutability and evolvability

# Indirect selection for mutability

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How readily does mutability evolve? Petrie and Roberts (2007) have recently described a theoretical example of increased mutation rate based on female choice. Mutator alleles can also be favored by strong selection for phenotypic variation, such as that imposed by immunological attack against pathogens, together with stable linkage to beneficial mutations, provided by haploidy in microorganisms. But the special conditions required for these examples highlight two assumptions that have framed discussion of mutation-rate evolution for most of the past century (for example, Bataillon, 2000; Bell, 2005; Cotton and Pomiankowski, 2007). First, although close linkage may allow a mutator to hitchhike on selection for a beneficial allele, recombination, at least in sexually reproducing populations, will eventually separate the two. Second, because most non-neutral mutations are deleterious, the net effect of any mutator must be fitness reduction. Thus, 'natural selection of mutation rates has only one possible direction, that of reducing the frequency of mutation to zero' (Williams, 1966). Regrettably, this classic but overstated conclusion remains influential. Even well-established exceptions like the 'contingency loci' of some bacteria are routinely marginalized as special cases that depend on extreme and/or unusual circumstances (Sniegowski and Murphy, 2006).

Contemporary discussion also often oversimplifies 'mutation rate' as a single statistic rather than a cumulative total emerging from many distinct mutational mechanisms (for example, Bataillon, 2000; Sniegowski and Murphy, 2006). However, although some mutator alleles may affect genome-wide accuracy of DNA processing, others have effects that are restricted to individual sites. Each site-specific mechanism may carry its own distinct rate and its own unique probability distribution for deleterious and beneficial effects. Such characteristics allow certain common mutagenic patterns to escape the reach of Williams' conclusion.

This is most clearly illustrated by the properties of simple sequence repeats

(SSRs, also termed microsatellites and minisatellites). SSR 'slippage' mutations, which increase or decrease the number of tandem repeats, occur at rates which may be orders of magnitude greater than those for single nucleotide substitutions. The particular mutation rate at each SSR depends on locus characteristics including motif length and purity of repetition. Because the rate-determining locus and the locus at which mutations occur are one and the same, there is no possibility for recombination to separate the two. Thus, an SSR locus represents a 'mutator allele' whose site for mutation is itself. Consequently, even in diploid, sexually reproducing genomes, an SSR mutator allele will always remain linked with its resulting mutations.

Although SSRs are often considered 'junk', SSR repeat-number alleles can influence almost any aspect of genetic function from protein coding to exon splicing to regulatory interaction. Quantitative functional effects have been reported for SSRs located in exons, introns, and upstream and downstream regulatory domains (Kashi and King, 2006). Although harmful SSR mutations do exist (for example, triplet repeat diseases), SSRs more typically yield 'mutations of small effect'. And 'mutations of small effect' are potentially beneficial with probability approaching 50% (Fisher, 1930). Hence the genetic variation supplied by SSR mutator loci need not be predominantly deleterious.

At any given SSR locus, each allele encodes both a phenotypic effect, represented by the number of repeats, and a mutation rate, represented by purity of motif repetition, for example (Trifonov, 1989). Therefore, natural selection acting on the fitness effects of SSR alleles also indirectly selects their mutation rates. Eliminating a high-mutation-rate allele proceeds gradually through repeated rounds of selection against individual deleterious mutants as they arise. But if a high-mutation-rate allele gives rise to a beneficial mutant, selection that fixes the mutant will unavoidably fix the high mutation rate as well. Thus, indirect selection can readily exploit the characteristic mutability of SSRs to minimize mutation rates at sites where variation is

disadvantageous while assuring that variability remains present at sites that repeatedly experience directional selection.

A number of observations indicate that SSRs are distributed non-randomly with respect to gene function (Kashi and King, 2006). A surprisingly large proportion of genes are closely associated with one or more SSRs, with especially high prevalence in regulatory loci. Triplet repeats are most common in protein-coding domains where they allow adjustment of length of amino-acid repeats. Non-triplet motifs predominate in other functional domains. The overall distribution of SSR motifs also varies among taxa, while SSRs in homologous locations may have different motifs in related species. Such patterns are readily interpretable as resulting from indirect selection for the site-specific mutability and allelic variation that SSRs provide.

Two common objections to the hypothesis of selection favoring mutability stem from an unfortunate conflation of 'mutability' with 'evolvability'. First, since individual organisms do not evolve, a population-level property like evolvability can be favored only by some form of group selection (Williams, 1966). Thus, the widely accepted implausibility of group selection also impugns selection for mutability (Sniegowski and Murphy, 2006). Second, selection for evolvability is often challenged simply because the advantages of future adaptation cannot be a selective force in the present (Sniegowski and Murphy, 2006). But in situations where variability offers immediate benefits, indirect selection for site-specific mutability, proceeding at the level of individual genes, is no less plausible than direct selection for fitness. Evolvability emerges as an epiphenomenon at the level of populations.

Williams (1966) wisely recognized that 'our current picture of evolutionary adaptation is, at best, oversimplified and naive'. Special conditions are certainly required before selection can favor mutability. But 'special' does not necessarily imply 'unusual'. SSRs illustrate just how readily appropriate conditions of site-specific mutability can obtain. Additional sources for genetic variation, such as transposable elements (for example, Capy *et al.*, 2000), may also be amenable to indirect selection. Just as sexual recombination offers advantageous shuffling of preexisting variation, so too may new variations, if suitably constrained by site-specific mechanisms, accrue substantial advantage. As Darwin recorded, 'some authors

believe it to be as much the function of the reproductive system to produce individual differences...as to make the child like its parents'. This point of view may be more pertinent to understanding the dynamic genome than Sturtevant's dismissive dictum that 'mutations are accidents, and accidents will happen'.

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- Bataillon T (2000). Estimation of spontaneous genome-wide mutation rate parameters: whither beneficial mutations? *Heredity* **84**: 497–501.
- Bell G (2005). The evolution of evolution. *Heredity* **94**: 1–2.

- Capy P, Gasperi G, Biéumont C, Bazin C (2000). Stress and transposable elements: co-evolution or useful parasites? *Heredity* **85**: 101–106.
- Cotton S, Pomiankowski A (2007). Sexually selected mutation rates. *Heredity* **98**: 185–186.
- Fisher RA (1930). *The Genetical Theory of Natural Selection*. Oxford University Press: Oxford.
- Kashi Y, King DG (2006). Simple sequence repeats as advantageous mutators in evolution. *Trends Genet* **22**: 253–259.
- Petrie M, Roberts G (2007). Sexual selection and the evolution of evolvability. *Heredity* **98**: 198–205.
- Sniegowski PD, Murphy HA (2006). Evolvability. *Curr Biol* **16**: R831–R834.
- Trifonov EN (1989). The multiple codes of nucleotide sequences. *Bull Math Biol* **51**: 417–432.
- Williams G (1966). *Adaptation and Natural Selection*. Princeton University Press: Princeton.

### Editor's suggested reading

- Caporale LH (2003). Natural selection and the emergence of a mutation phenotype: an update

- of the evolutionary synthesis considering mechanisms that affect genomic variation. *Annu Rev Microbiol* **57**: 465–485.
- Fondon III JW, Garner HR (2004). Molecular origins of rapid and continuous morphological evolution. *Proc Natl Acad Sci USA* **101**: 18058–18063.
- Kirschner M, Gerhart J (1998). Evolvability. *Proc Natl Acad Sci USA* **95**: 8420–8427.
- Li Y-C, Korol AB, Fahima T, Beiles A, Nevo E (2002). Microsatellites: genomic distribution, putative functions and mutational mechanisms: a review. *Mol Ecol* **11**: 2453–2465.
- Moxon ER, Thaler DS (1997). The tinkerer's evolving tool-box. *Nature* **387**: 659–662.
- Theodorou K, Couvet D (2006). Genetic load in subdivided populations: interactions between the migration rate, the size and the number of subpopulations. *Heredity* **96**: 69–78.
- Worobey M (2005). Genomics: anthrax and the art of war (against ascertainment bias). *Heredity* **94**: 459–460.