

Evolutionary Genomics

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Supplement Aims and Scope

This supplement is intended to focus on evolutionary genomics. *Evolutionary Bioinformatics* aims to provide researchers working in this complex, quickly developing field with online, open access to highly relevant scholarly articles by leading international researchers. In a field where the literature is ever-expanding, researchers increasingly need access to up-to-date, high quality

D volutionary genomics covers a wide range of subjects investigating the evolution of species' genomes.¹ This supplement presents various models of evolution, from bacteria² and plants³ to humans.^{4–7} Indeed, the similarities in genetic codes across all types of organisms allow for comparisons of DNA sequences between and within species.⁸ Interpretation of these comparisons can nonetheless be challenging because of the diversity of approaches available and the way genotypes are determined, ie genotyping or sequencing techniques used, and characteristics of the markers or loci of interest.⁵

The extent to which observed changes in DNA sequences are adaptive has always been central to the understanding of evolution and is reflected here by the number of papers addressing the effect of natural selection on observed patterns of diversity.^{3,4,8} Furthermore, other processes, including genetic drift, gene overlap, migration, mutation and recombination interact with selection to shape patterns of diversity.⁶ These interactions can lead to counterintuitive outcomes. For example, the highest frequencies of the allele associated with lactase persistence in Europe are found in the North-West. This scholarly articles on areas of specific contemporary interest. This supplement aims to address this by presenting high-quality articles that allow readers to distinguish the signal from the noise. The editor in chief hopes that through this effort, practitioners and researchers will be aided in finding answers to some of the most complex and pressing issues of our time.

may lead to the assumption that the allele arose and diffused from there. However, computer simulations have shown that the joint effect of positive selection and allele-surfing with the spread of farmers from the Near East about 8,000 years ago could lead to a similar pattern.^{9,10} Computational models continue to be insightful analytic tools to understand the interplay between demographic and adaptation processes.^{3,4,9}

Recent advances in high throughput genomics techniques provide better coverage of organism populations, and genomes. Consequently, the field of evolutionary genetics that was mainly model-based in the 20th century has more recently been driven by big data to become the relatively new field of evolutionary genomics. Genome-wide surveys of populations have proven power to detect associations between genetic variants and diseases or complex traits and to investigate for the effects of selection.¹¹ They show that evolutionary genetics is more relevant to practical problems than ever before, such as plant improvement³ and human health,⁷ where evolutionary insights aid in the identification of patients' disease-causing alleles and genes.^{12,13} A single change in the DNA can have multiple consequences at the organism level.¹⁴ Studies that will approach these various aspects of a change in the DNA, including genomic context, functional effect, and evolutionary modeling (ways an adaptive change increases in frequency in a population), will provide a better understanding of the adaptability of organisms and species.¹⁵

Evolutionary genomics could also be applied to identify potentially beneficial genetic variants from the single gene to the whole genome scale, for human needs such as for the production of commodity chemicals and pharmaceuticals. Such genomic approaches are mainly directed at two extremes: (1) focusing on a single or few genomic loci to thoroughly investigate responses to localized edits, such as the recombineering technique used mainly in E. coli;^{16,17} and (2) an approach termed Adaptive Laboratory Evolution (ALE) in which cells are being challenged with an environmental stress and adapt to regain fitness.^{18,19} Between these two extremes, one using prior knowledge and rational design and the other a completely unbiased ALE approach, other approaches emerged that try to incorporate the best of each method. Such approaches, such as MAGE and TRMR,^{20,21} combine rational design with multiple genome-wide edits. Recent advances in genome engineering tools such as the CRISPR-CAS9 system^{22,23} hold promise for even more precise and systematic editing technologies, which will allow rational whole genome surveys for specific traits at the single nucleotide resolution.

Future directions in evolutionary genomics may apply biological distance metrics alternative to sequence similarity, such as the human gene connectome,^{24,25} gene-level quantitative accumulated mutational damage and intolerance, and gene-specific mutation impact predictions to complement traditional selective pressure approaches.^{26–28} In the era of high throughput data, efficient computational methodologies are essential.²⁹ A strong interdisciplinary effort will facilitate solving the various challenges in today's evolutionary genomics.

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