Spatial Patterns of Molecular Trends in Bacterial Genomes

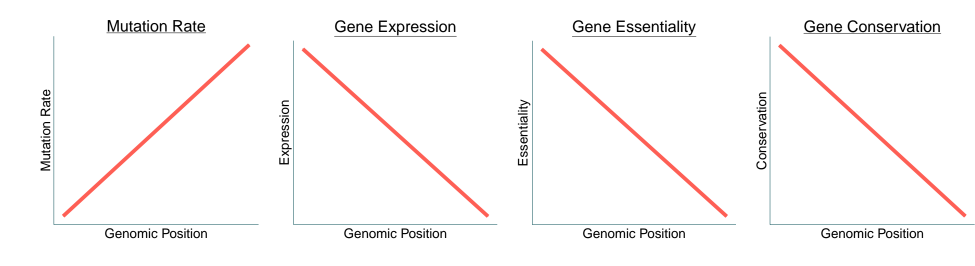
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Molecular Traits Vary with Genomic Position

Molecular traits such as mutation rate and gene expression change with distance from the origin of replication. Genes that are highly conserved, essential, and expressed at higher levels are typically found near the origin of replication $^{1;2;3;4;5}$. The disparity in molecular features between the origin of replication and the terminus is thought to be due to decreased mutation rate near the origin of replication 1;3;4 and the requirement to maintain proper gene dosage³ amount and replication timing $^{1;5}$. These **spatial** molecular trends have not been analyzed while accounting for genomic reorganization such as rearrangements and inversion, which provide bacteria with the opportunity to gain new genetic information.



Objective: In depth analysis of genomic patterns of substitutions and gene expression in bacterial genomes.

Hypothesis: The number of substitutions increases with increasing distance from the origin of replication.

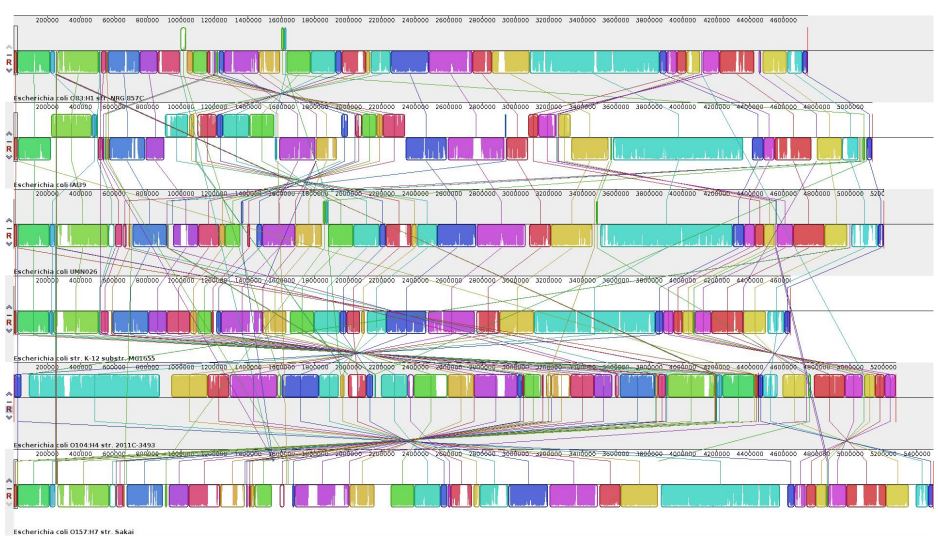
1. Methods

25 bacterial genomes and gene expression data from 9 bacterial species.

Bacteria and Replicon	Number of Strains	Genome Structure
Escherichia coli Chromosome	6	0
Bacillus subtilis Chromosome	7	0
Streptomyces Chromosome	6	
Sinorhizobium meliloti Chromosome	6	\bigcirc \circ \circ
Sinorhizobium meliloti pSymA	6	$\bigcirc \circ \circ$
Sinorhizobium meliloti pSymB	6	$\bigcirc \circ \bigcirc$

Alignment:

- 1. Each whole genome was **globally aligned** to obtain **L**ocally **C**o-linear Blocks (LCB) which allows for genome reorganization such as inversions and rearrangements (progressiveMauve⁶).
- 2. **Each LCB** was **locally re-aligned** with MAFFT⁷ for a more accurate alignment.



Example progressiveMauve alignment of *E. coli* genomes.

Phylogeny:

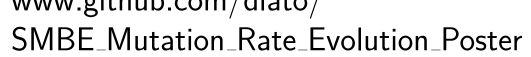
3. Bootstrap phylogenetic trees were constructed for each bacterial replicon (PHYLIP⁸).

Statistical Analysis:

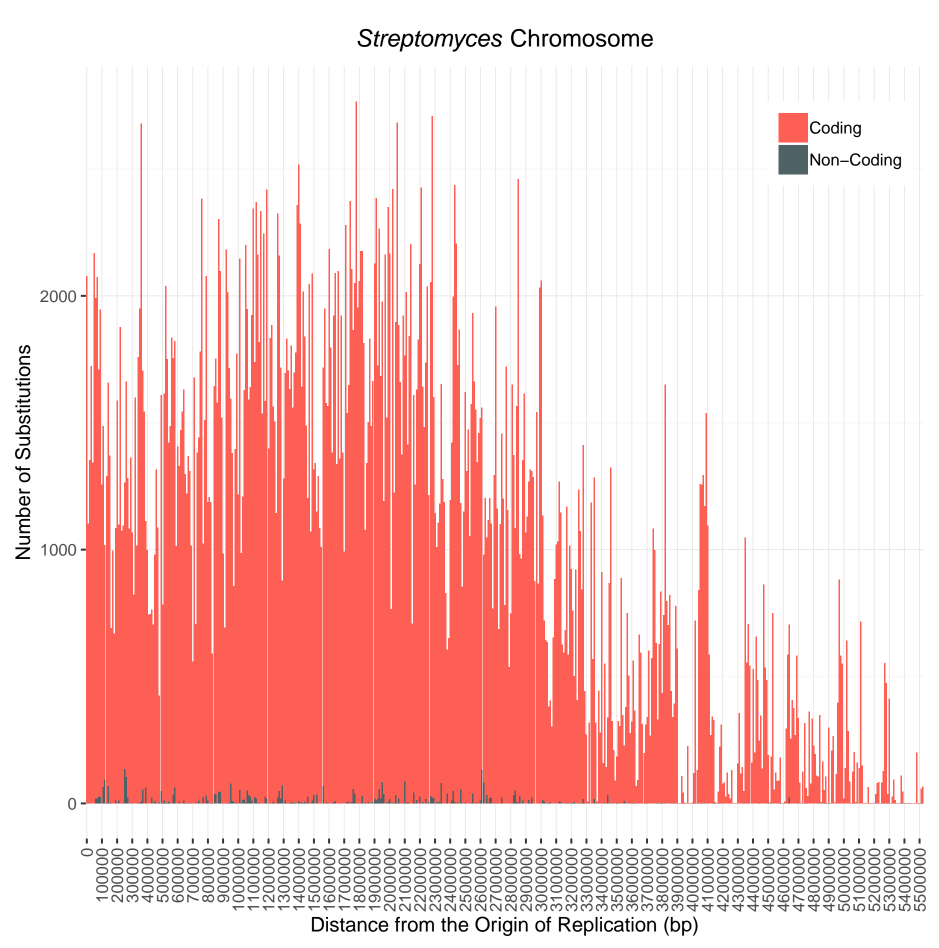
- 4. Each **genomic position** was **scaled** to the origin of replication and accounted for bidirectional replication (R^9) .
- 5. To track genome rearrangements, nucleotide substitutions and genomic positions were reconstructed in extinct taxa and ancestors on the given phylogenies (custom Perl and Python scripts, $PAML^{10}$).
- 6. Logistic regression and linear regression were performed to compute changes in substitutions and gene expression across all coding and non-coding segments of the genome (R^9) .

See GitHub for more information.

www.github.com/dlato/



2. Substitutions \ with Genomic Position



Bacteria and Replicon	Coefficient Estimate		Genome Structure
	Coding Sites	Non-Coding Sites	
E. coli Chromosome	-9.119×10 ⁻⁸ ***		0
B. subtilis Chromosome	-1.273×10 ⁻⁷ ***	-9.861×10 ⁸ ***	0
Streptomyces Chromosome			
S. meliloti Chromosome	$-1.550 \times 10^{-7} ***$	-1.510×10^{-7} *	000
S. meliloti pSymA	-1.156×10^{-7} *	NS	000
S. meliloti pSymB	$2.587 \times 10^{-7} ***$	$8.591 \times 10^{-7} ***$	\circ

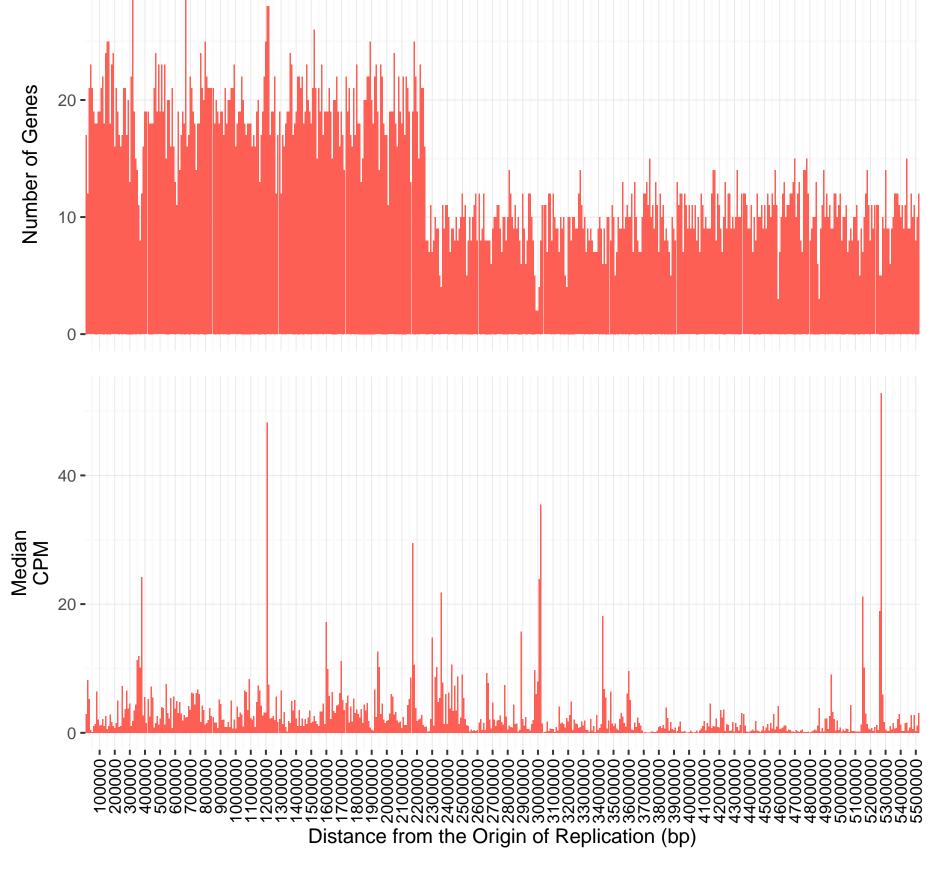
Significance Codes: p < 0.001 = '***', 0.001 < 0.01 = '**', 0.01 < 0.05 = '*', NS =Not Significant

Discussion: Increased transposon insertion events 11 and potential genomic and pathogenicity islands $^{12;13}$ near the origin of replication can cause asymmetry in nucleotide composition 14 , GC content 15 , and mutation rate $^{1;3;4}$. This could be why we see an increase in the number of substitutions near the origin of replication.

Conclusion: The number of substitutions decreases with increasing distance from the origin of replication in most bacterial replicons at most coding sites.

3. Gene Expression \ with Genomic Position

Streptomyces Chromosome



Coefficient Estimate	Genome Structure
-6.03×10 ⁻⁵ ***	0
	0
-1.17×10 ⁻⁶ ***	
NS	000
1.39×10^{-3} ***	$\bigcirc \circ \circ$
NS	$\bigcirc \circ \circ$
	$-6.03 \times 10^{-5}***$ $-9.7 \times 10^{-5}***$ $-1.17 \times 10^{-6}***$ NS $1.39 \times 10^{-3}***$

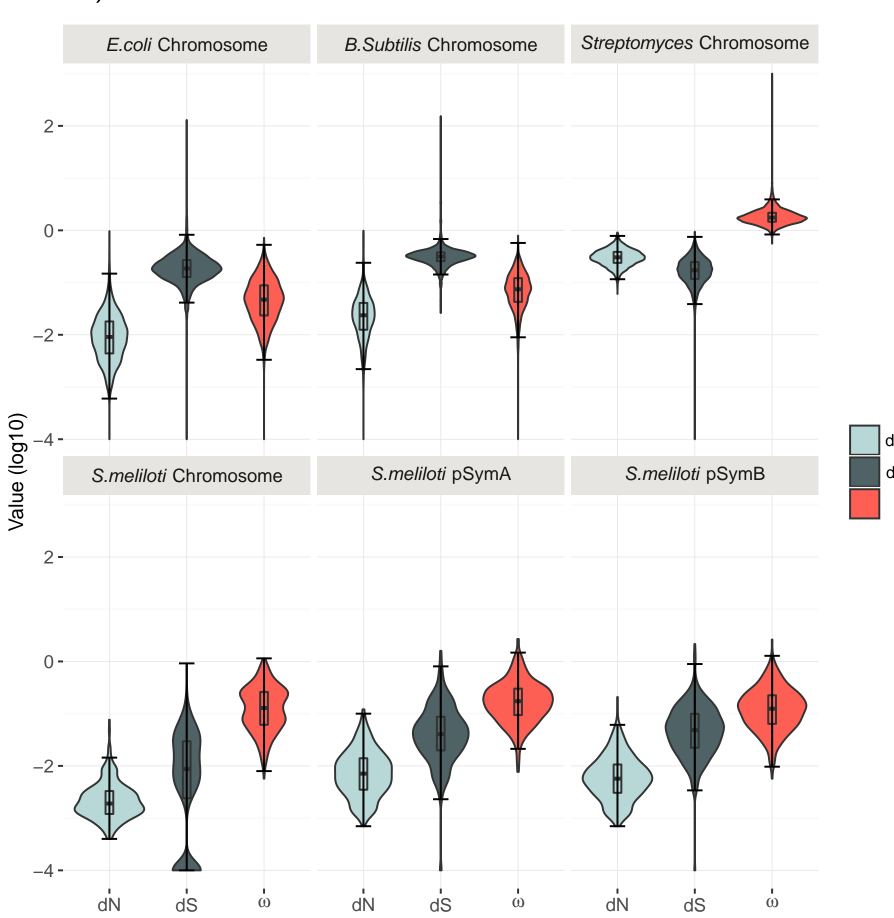
Significance Codes: p < 0.001 = "**", 0.001 < 0.01 = "*", 0.01 < 0.05 = "*", NS = ""."Not Significant

Discussion: Replication error increases when moving away from the origin of replication, therefore, genes that are highly expressed are often located near the origin. 16;17;18

Conclusion: Gene expression decreases with increasing distance from the origin of replication in most bacterial replicons.

4. Preliminary Selection Results

Distribution of dN, dS, and ω for the coding regions (codeml $PAML^{10}$).



dN = # of non-synonymous substitutions per site, dS = # of synonymous substitutions per site, $\omega = \mathsf{ratio} \; dN/ \; dS$

Most of the bacterial replicons are under **purifying selection** ($\omega < 1$).

Conclusions and Ongoing Research

Determining how the number of substitutions are distributed spatially throughout bacterial genomes broadens our understanding of their evolution. Most coding regions of the replicons considered in this study have the number of substitutions and gene expression decrease with increasing distance from the origin of replication. The exceptions to this were pSymB of S. meliloti, E. coli and Streptomyces which had the number of substitutions increase when moving away from the origin of replication. These spatial substitution and gene expression results can be used to determine if all bacteria possess the same evolutionary patterns.

Current work involves:

- Determining the dN/dS ratio for each coding section of the bacterial genomes and if this changes with genomic position.
- Identifying inversions within bacterial genomes and how they might alter gene expression
- Performing ancestral reconstruction of gene expression in several *E. coli* species

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