Subs Paper Things to Do:

- why are the lin reg of dN, dS and ω NS but the subs graphs are...explain!
- mol clock for my analysis?
- GC content? COG? where do these fit?

Inversions and Gene Expression Letter Things to Do:

- create latex template for paper
- look up inversions and small RNA's paper Marie was talking about at Committee meeting
- write outline for letter
- write Abstract
- write intro
- write methods
- compile tables (supplementary)
- write results
- write discussion
- write conclusion
- do same ancestral/phylogenetic analysis that I did in the subs paper

General Things to Do:

• summarize references 40 and 56 from Committee meeting report (Brian was asking)

Last Week

Substitutions Paper:

✓ finished latest draft of second round of revisions

✓submit!

Inversions + Gene Expression:

✓ finished first draft of inversions paper (minus the DESeq analysis)

- ✓ looked at some HGT genes and inversions
- ✓ phylogentic tree for the taxa
- ✓ HNS and expression values for bound and unbound regions

Dissertation:

- ✓ Finished inversions proposed studies
- ✓ Finished inversions conclusion thesis summary
- ✓ Started inversions discussion

Inversions + Gene Expression: I attempted to use the individual sample names as a component in the DESeq analysis (sample_name ~ treatment). But it did not work. DESeq still complains that the matrix is not full rank and the column factors (sample_name and treatment) are linear combinations of each other. I do not know what to do. I need to account for variation in expression between datasets (since they were all done in different labs at different times), while still being able to see what differences are due to the "treatment" = inversions. Do you know how I can achieve this?

I realized that I was considering a "match" between HNS and my inversions if any part of an HNS binding site overlapped with any region within the entire inverted block. This meant that each gene within an inverted block would be considered "bound" to HNS. I think this is wrong, so I instead considered a "match" to be if any part of the HNS binding site overlapped with any portion of a gene. This way each individual gene would be assessed as to if it had HNS bound or not. Figure 2 and Table 4 are updated with this information. There is now complete consistency in sign between the significant correlations (Table 4). However, I am not sure which genomes to use for the HNS binding. The data for HNS binding sites is based on the *E. coli* K-12 MG1655 strain. So I think that I should only be looking at this genome. However, the *E. coli* K-12 MG1655 strain has no inverions, they mostly occur in the *E. coli* ATCC strain. So I am considering an inverted alignment block as having at least one taxa inverted. Do you think it is ok that I am only looking at *E. coli* K-12 MG1655 and using the above criteria for inverted blocks to see if there is a correlation between HNS and inversions?

Since a lot of the genes that HNS represses are linked to being recently horizontally transferred, I obtained the HGT information from the significant HNS datasets which is from previously published studies on HGT. I looked at the correlation (Pearson) between HGT genes and inverted sequences, and HGT and blocks with significant differences in gene expression and I found no significant correlation for either. Should I be pulling HGT data from other sources to look into this? I just thought it would be good to at least touch on this in the paper since HGT genes are mentioned in a lot of the HNS literature.

This Week

- get HGT info for Oshima HNS data
- do HGT correlations on ↑ data
- Dissertation: discussion on inversions paper
- Dissertation: intro for inversions paper
- actual analysis on DESeq data
- visualizations/results for \ \
- start working on exit seminar presentation

Next Week

- continue working on exit seminar presentation
- go over missing spots/red marks in conclusion
- edit conclusion
- Brian's edits to inversions paper
- maybe do inversions in 10kb blocks? (and other sliding windows?)
- dist from ori on DESeq results?

Bacteria and Replicon	Protein Coding Sequences Coefficient Estimate
E. coli Chromosome	$-3.29 \times 10^{-8***}$
$B.\ subtilis\ { m Chromosome}$	8.70×10^{-9} *
Streptomyces Chromosome	NS
$S.\ meliloti\ { m Chromosome}$	$-6.80 \times 10^{-7***}$
S. meliloti pSymA	$4.49 \times 10^{-7***}$
$S.\ meliloti\ pSymB$	6.27×10^{-8} *

Table 1: Logistic regression analysis of the number of substitutions along all protein coding positions of the genome of the respective bacteria replicons. ONLY EXTANT BRANCHES. Grey coloured boxes indicate a negative logistic regression coefficient estimate. All results are statistically significant. Logistic regression was calculated after the origin of replication was moved to the beginning of the genome and all subsequent positions were scaled around the origin accounting for bidirectional replication. All results are marked with significance codes as followed: < 0.001 = '***, 0.001 < 0.01 = '**, 0.001 < 0.01 = '**, 0.001 < 0.05 = '*, 0.001 < 0.05 = 'NS'.

Average Gene Expression within Alignment Blocks • Non-significant—Inverted • Non-inverted

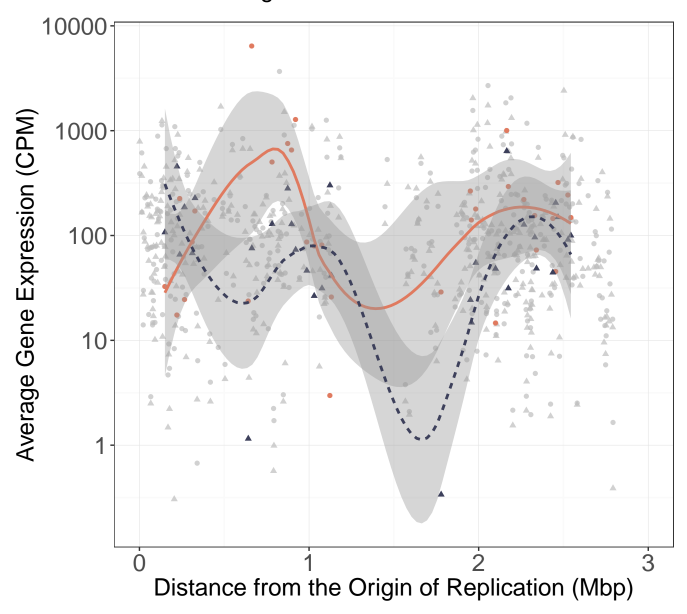


Figure 1: Visualization of the difference in gene expression between inverted and non-inverted sequences within alignment blocks. Each alignment block represents homologous sequences between the Escherichia coli strains insert table ref here. E. coli K-12 MG1655 was used as the reference genome for genomic position for each alignment block. The midpoint of each alignment block was calculated to be the genomic distance from the E. coli K-12 MG1655 origin of replication. Each alignment block has one point on the graph to represent the average expression value in Counts Per Million (CPM) for all inverted (circles) and non-inverted (triangles) sequences within the block. Blocks that had a significant difference in gene expression (using a Wilcoxon sign-ranked test, see Materials and Methods) have the inverted and non-inverted gene expression averages highlighted in pink circles and purple triangles respectively. A smoothing line (loewss) was added to link the average gene expression values for the inverted (pink solid) and non-inverted (purple dashed) sequences within block that had a significant difference in gene expression (using a Wilcoxon signranked test, see Materials and Methods). All blocks that did not have a significant difference in average gene expression between inverted and non-inverted sequences within alignment blocks have the average inversion (circles) and non-inversion (triangles) gene expression values coloured in light grey.

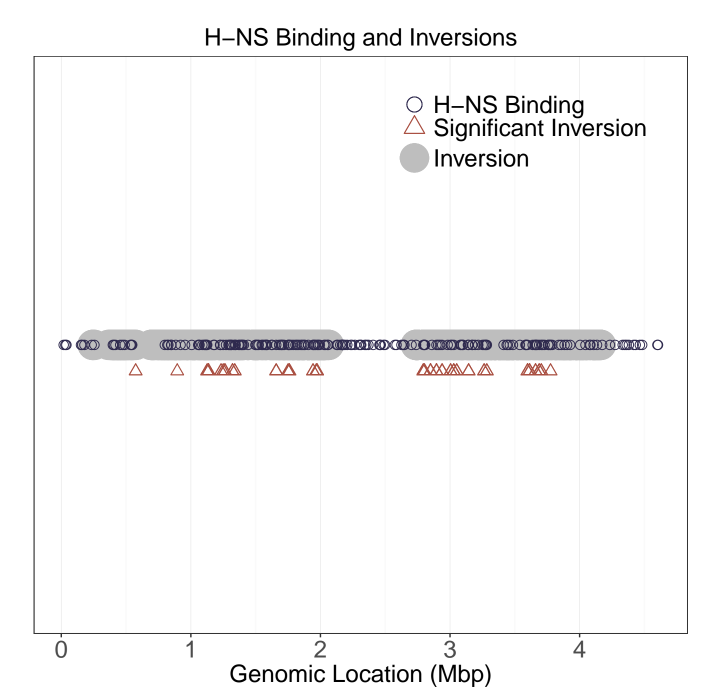


Figure 2: Visualization of the genomic locations of all inversion alignment blocks (light grey filled circles) identified between E. coli K-12 MG1655, E. coli K-12 DH10B, E. coli BW25113, and E. coli ATCC. The data points are plotted on the genome of E. coli K-12 MG1655 which is used as a reference. Each inversion alignment block has a single genomic location chosen to be the midpoint of the inverted region calculated to be the genomic distance from the E. coli K-12 MG1655 origin of replication. Histone-like Nucleoid-Structuring (H-NS) protein binding sites in the E. coli K-12 MG1655 are overlaid on top of the inversion alignment blocks (circles outlined in dark purple). Data for the H-NS binding information is from Higashi insert citation here. Inversion alignment blocks that had a significant difference in gene expression between the inverted and non-inverted sequences within the block (using a Wilcoxon sign-ranked test, see Materials and Methods), are marked below the inverted alignment blocks with dark pink outlined triangles.

Strain Removed	Coefficient Estimate		
	$E.\ coli$		
None	$-2.66 \times 10^{-8***}$		
U00096	$-3.12 \times 10^{-8***}$		
CP0032890	$-3.07 \times 10^{-8***}$		
CU9281640	$-2.95 \times 10^{-8***}$		
CP0018550	$-1.50 \times 10^{-8***}$		
BA0000070	$-2.63 \times 10^{-8***}$		
CU9281630	$-2.49 \times 10^{-8***}$		
$B.\ subtilis$			
None	$2.76 \times 10^{-8***}$		
NC_000964	$2.96 \times 10^{-8***}$		
NC_018520	$3.57 \times 10^{-8***}$		
NC_017195	$1.00 \times 10^{-7***}$		
NC_022898	$5.17 \times 10^{-8} ***$		
NC_014976	$-4.02 \times 10^{-8***}$		
CP01731	$5.43 \times 10^{-8***}$		
NC_014479	NS		
Streptomyces			
None	$7.21 \times 10^{-8***}$		
CP050522	$8.37 \times 10^{-8***}$		
GG657756	$3.62 \times 10^{-8***}$		
CP042324	$7.72 \times 10^{-8***}$		
AL645882	$7.65 \times 10^{-8***}$		
CM001889	$-2.46 \times 10^{-7***}$		

Table 2: Logistic regression on the presence or absence of a substitution and distance from the origin of replication. Each strain was systematically removed and the entire analysis was repeated. All results are marked with significance codes as followed: <0.001= '***', 0.001<0.01= '**', 0.01<0.05= 'NS'.

Strain Removed	Coefficient Estimate	
S. meliloti Chromosome		
None	$-6.57 \times 10^{-7***}$	
NC_015590	$-3.18 \times 10^{-7***}$	
NC_003047	$-6.01 \times 10^{-7***}$	
CP004140	$-6.00 \times 10^{-7***}$	
CP009144	$-6.67 \times 10^{-7***}$	
NC_017322	$-7.19 \times 10^{-7***}$	
NC_017325	$-5.01 \times 10^{-7***}$	
$S.\ melil$	oti pSymA	
None	$2.74 \times 10^{-7} ***$	
NC_017327	$6.98 \times 10^{-7***}$	
CP009145	$1.78 \times 10^{-7***}$	
NC_003037	$2.09 \times 10^{-7***}$	
CP004138	$2.08 \times 10^{-7} ***$	
NC_015591	NS	
NC_017324	$-1.52 \times 10^{-6***}$	
S. melil	oti pSymB	
None	$1.10 \times 10^{-7***}$	
NC_015596	$6.78 \times 10^{-7***}$	
NC_017326	$1.67 \times 10^{-7} ***$	
NC_017323	NS	
CP009146	$-2.57 \times 10^{-7***}$	
CP004139	$1.04 \times 10^{-7***}$	
NC_{003078}	$1.04 \times 10^{-7***}$	

Table 3: Logistic regression on the presence or absence of a substitution and distance from the origin of replication. Each strain was systematically removed and the entire analysis was repeated. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "*", 0.01 < 0.05 = "", 0.05 = "NS".

H-NS Binding Study	All Inversions H-NS Binding	Significant Inversions and H-NS Binding	Total Number of H-NS Binding Sites Within All Alignment Blocks
Grainger 2006 [?]	NS	NS	53
Ueda 2013 [?]	NS	NS	275
Higashi 2016 [?]			
criteria A	0.0467*	NS	371
criteria B	0.0540**	NS	343
criteria C	0.0540**	NS	343
criteria D	0.0540**	NS	343
criteria E	0.0544**	NS	340
criteria F	0.0544**	NS	340
Lang 2007 [?]	0.0574**	NS	115
Oshima 2006 [?]	0.0390*	NS	664

Table 4: are there any other stats related to correlation that people like to have in these tables that I should also be including? Pearson correlation between H-NS binding sites and inverted regions of the E. coli K-12 MG1655 genome. A genomic region was considered inverted if this sequence was inverted in any of the following four taxa: E. coli K-12 MG1655, E. coli K-12 DH10B, E. coli BW25113, and E. coli ATCC. The genomic positions of these inversions in E. coli K-12 MG1655 was used for reference. The binding sites for the H-NS protein are in the genomic coordinates of E. coli K-12 MG1655, chosen as a reference. The second column "All Inversions and H-NS Binding" represents the correlation coefficient between inverted regions and H-NS binding sites. The third column "Significant Inversions and H-NS Binding" represents the correlation coefficient between inverted regions with significant differences in normalized gene expression between inverted and non-inverted taxa (via a Wilcoxon signed-rank test) and H-NS binding sites. The ref Higashi data set had multiple criteria to define H-NS binding sites. They are listed as follows: A: Genes whose coding regions overlap with the H-NS binding regions, B: Genes whose coding regions overlap with the H-NS binding regions and intergenic regions that were bound by H-NS, C: Genes whose coding regions overlap with the H-NS binding regions and intergenic regions that are "class I " (see cite Higashi), D: Genes whose coding regions overlap with the H-NS binding regions and intergenic regions that contain known promoter sequences, E: Same as A, but genes on which H-NS binding is restricted to the 3' end and the length overlapping with H-NS-bound regions is <10\% of the total gene length were excluded from H-NS-bound genes, F: When genes included in transcriptional units whose upstream regions or first coding regions overlapped with H-NS bound regions, all genes in the transcriptional units were judged as genes affected by H-NS binding. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "**", 0.01 < 0.05 = "*", > 0.05= 'NS'.

Datasets:	Correlation Coefficient (W	
Inverted Blocks	15218699**	
Inverted Sequences	11436344***	

Table 5: Correlation coefficients for Wilcoxon signed-rank test on various datasets to determine the correlation between an inversion and difference in normalized gene expression. The "Inverted Blocks" dataset represents alignment blocks that have at least one taxa with an inverted sequence. The "Inverted Sequences" dataset represents all individual sequences from all alignment blocks that were inverted. The correlation between both datasets was computed using a Wilcoxon signed-rank test. All results are marked with significance codes as followed: < 0.001 ="**", 0.001 < 0.05 = "*", > 0.05 = "NS".

% of Blocks that are		
Inverted	Inverted with	Increased in
	Differences in	Gene Expression
Gene Expression in Inverted		in Inverted Sequences
68.29	8.22	58.06

Table 6: Percent of blocks in categories for various datasets (blocks with all 4 taxa, at least 3 taxa, or at least 2 taxa). The second column is any block that had at least one sequences that was inverted. The last column only deals with blocks that had at least one inverted sequence and had a significant difference in gene expression (column 3).

Block Length Correlation Coefficient (W)

4060729.5***

Table 7: Correlation coefficients for Wilcoxon signed-rank test in alignment blocks. The correlation coefficient represents a correlation between alignment block length and blocks with a significant/non-significant difference in normalized gene expression between inverted and non-inverted sequences within the block. All results are marked with significance codes as followed: < 0.001 = '**', 0.001 = '**', 0.01 < 0.05 = '*', > 0.05 = 'NS'.

Genomic Position Correlation Coefficient (W)

NS

Table 8: Correlation coefficients for Wilcoxon signed-rank test in alignment blocks with a significant difference in normalized gene expression between inverted and non-inverted sequences within the block. The correlation coefficient between the significant blocks and the genomic position of the alignment blocks. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 = "*", 0.01 < 0.05 = "*", 0.005 = "NS".

Inversion Category	Correlation Coefficient
rev comp	NS
inversion	$2.20 \times 10^{-7} ***$
sig rev comp	-1.89×10^{-7} *
$sig \sim midpoint all blocks$	NS
$sig \sim midpoint inverted blocks$	NS

Table 9: Logistic regression between various inversion categories and distance from the origin of replication for all strains. rev comp = individual sequences inverted, inversion = block that has at least one inverted sequence, midpoint = block midpoint, sig = blocks with significant difference in normalized gene expression between inverted and non-inverted sequences within the block. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "**", 0.001 < 0.01 = "*", 0.001 < 0.05 = "NS".

Strain	rev comp	inversion
E. coli K-12 MG1655 E. coli K-12 DH10B	NS	$3.55 \times 10^{-7***}$ $3.45 \times 10^{-7***}$
E. coli BW25113	NS	$3.43 \times 10^{-7***}$ $3.73 \times 10^{-7***}$
E. coli ATCC	$-1.92 \times 10^{-7***}$	$-1.92 \times 10^{-7***}$

Table 10: Logistic regression between various inversion categories and distance from the origin of replication for each strain. rev comp = individual sequences inverted, inversion = block that has at least one inverted sequence, sig = blocks with significant difference in normalized gene expression between inverted and non-inverted sequences within the block. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "**", 0.01 < 0.05 = "", > 0.05 = "NS".