Subs Paper Things to Do:

- causes for weird selection and subs results in Streptomyces
 - see how often class 4 arises in strep to see what is going on in later portion of the genome (to see if annotation is really a problem)
 - split up the strep data into core and non core and see if results are the same
- make graphs proportional to length of respective cod/non-cod regions
- test examples for genes near and far from terminus (robust log reg/results)
- linear regression on 10kb regions for weighted and non-weighted substitutions
- average number of substitutions in 20kb regions near and far from the origin
- figure out why the data is weird for number of cod/non-cod sites
- why are the lin reg of dN, dS and ω NS but the subs graphs are...explain!
- grey out outliers in subs graphs?
- mol clock for my analysis?
- GC content? COG? where do these fit?

Gene Expression Paper Things to Do:

- linear regression on 10kb regions
- put new 10kb lin reg and # of genes over 10kb lin reg into paper
- write about \(\gamma\) in methods and discussion
- put expression lin reg and # coding sites log reg into supplement
- write about \(\gamma\) in paper and how results are the same
- update supplementary figures/file
- correlation of gene expression across strains
 - make graphs pretty and more informative with label names
 - add them to supplement with a mini write up of what we did and why
 - mention this in the actual paper
- if necessary add a phylogenetic component to the analysis
- potentially remove genes that have been recently translocated from the analysis
- model gene exp + position + number of genes

- split up the strep data into core and non core and see if results are the same
- what is going on with Streptomyces number of genes changing drastically from core to non-core
- codon bias?
- what is going on with really high gene expression bars
- edit paper
- submit paper

Inversions and Gene Expression Letter Things to Do:

- check if opposite strand in progressiveMauve means an inversions (check visual matches the xmfa)
- check if PARSNP and progressive Mauve both identify the same inversions (check xmfa file)
- create latex template for paper
- put notes from papers into doc
- use large PARSNP alignment to identify inversions
- confirm inversions with dot plot
- make dot plot of just gene presence and absence matrix (instead of each site) to see if this will go better
- 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)
- read and make notes on papers I found for dissertation intro

Last Week

✓ edits for gene expression intro

 \checkmark added some more references to the substitutions paper intro

✓ updated results from re-running substitutions analysis. (see tables below)

proportional coding subs > proportional non-coding subs

So I thought that I fixed this, which I did in pSymB, but when I re-ran the pipeline on the other bacteria the number of 10kb sections with proportionally more coding subs than non-coding subs still ranged from 14% - 50% of the 10kb sections. Looking into it a bit more, and it seems as though for the rest of the bacteria there is less data at a lot of the 10kb sections than there should be. So I need to figure out what is happening here.

dN/dS > 1 for some genes

Excluding Streptomyces, the rest of the bacteria have very few "genes" that have have dN/dS > 1 (4% of all "genes"). Looking into this further these "genes" are actually segments of genes that were really short (average of 33bp). This was because there were gaps in the genes so the rest of the gene got "cut" so it makes sense why these genes have such high ω . I am not sure if we should be discarding these gene fragments from the analysis? Thoughts?

Weird Streptomyces selection issue

I already talked to you about this, but I looked into why Streptomyces has so many genes with $\omega > 1$. The alignments are all correct and the genomic positions are all correct. So progressive Mauve and MAFFT are grabbing the correct genomic sequence. I BLASTed a coding gene from one of the blocks and it does match with sequences from all the other species of Streptomyces, however they are not found within the same block. My speculation is that progressive Mauve can not handle the slight differences between S. venezuelae and S. coelicolor and S. lividans and cuts the genome into too many blocks, classifying what genes are homologous. This can also be seen in the progressive Mauve block alignments where it looks like there is a clear divide between the sequence content of S. coelicolor and S. lividans and all the S. venezuelae strains. There is also very few sections (just by looking at the alignment) where all 6 taxa have sequence data and no gaps. Additionally, the branch lengths between the S. coelicolor and S. lividans strains are very small compared to the branch lengths between all the S. venezuelae strains. See below for what I plan to do next to fix all this.

Alignment split

I noticed that one of the blocks in *Streptomyces* failed to be split into coding and non-coding sections. I thought it was because I did not properly define an array v.s. a string for one of my variables, but I think something else is going on. Need to look into this more.

This Week

- look for more Streptomyces complete genomes to use in the analysis
- look into gblocks to assess MAFFT alignments
- check coding/non-coding code to see how often sites do not match up for Streptomyces
- check how often the tree for each block matches the overall tree for *Streptomyces*
- figure out what is going on with cod subs > non-cod subs
- deal with genes where $\omega > 1$

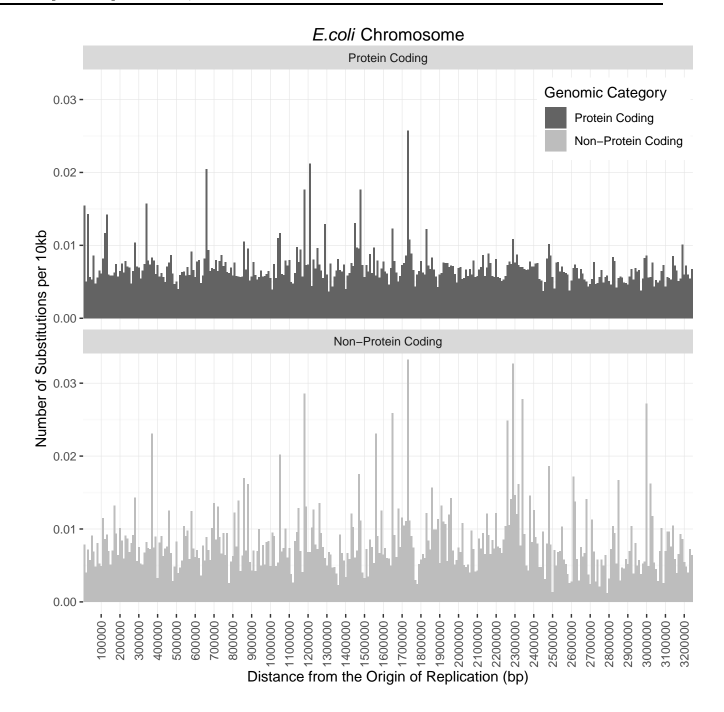
Next Week

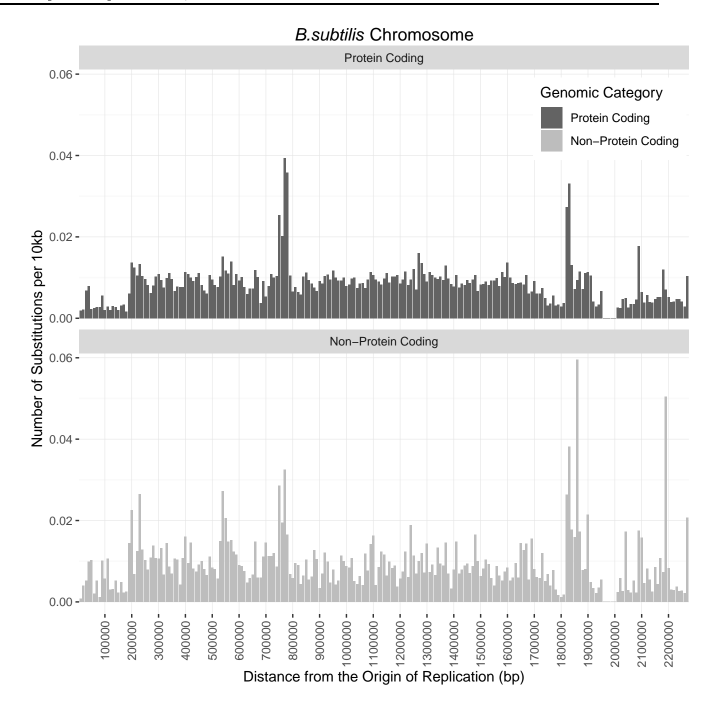
I would like to switch and work on the gene expression paper again.

- 1. phylogenetic analysis with gene expression in *E. coli*?
- 2. remove genes that have been recently translocated from analysis?
- 3. model gene expression + position + number of genes
- 4. split up *Streptomyces* data into core and non-core and see if the results are the same (do same for number of genes)

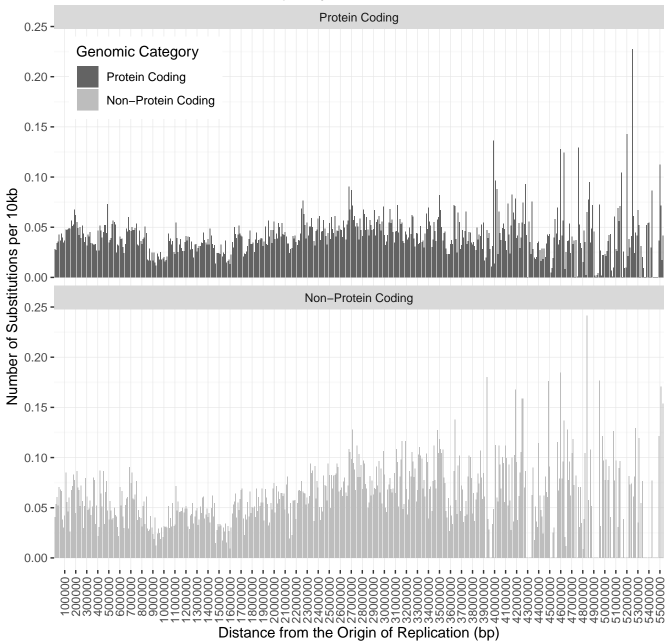
| | Near Origin | | | Near Terminus | | | |
|--------------------------------|-------------------------|----|-------------------------|---------------------------|--------------------------|----|--|
| Bacteria and Replicon | \overline{dN} | dS | ω | dN | dS | ω | |
| E. coli Chromosome | NS | NS | NS | NS | NS | NS | |
| B. subtilis Chromosome | NS | NS | NS | NS | NS | NS | |
| Streptomyces Chromosome | NS | NS | NS | NS | NS | NS | |
| $S.\ meliloti$ Chromosome | 2.79×10^{-8} * | NS | NS | NS | NS | NS | |
| $S.\ meliloti\ \mathrm{pSymA}$ | NS | NS | 3.42×10^{-5} * | NS | NS | NS | |
| $S.\ meliloti\ pSymB$ | NS | NS | NS | $-3.24 \times 10^{-7} **$ | $8.33 \times 10^{-6***}$ | NS | |

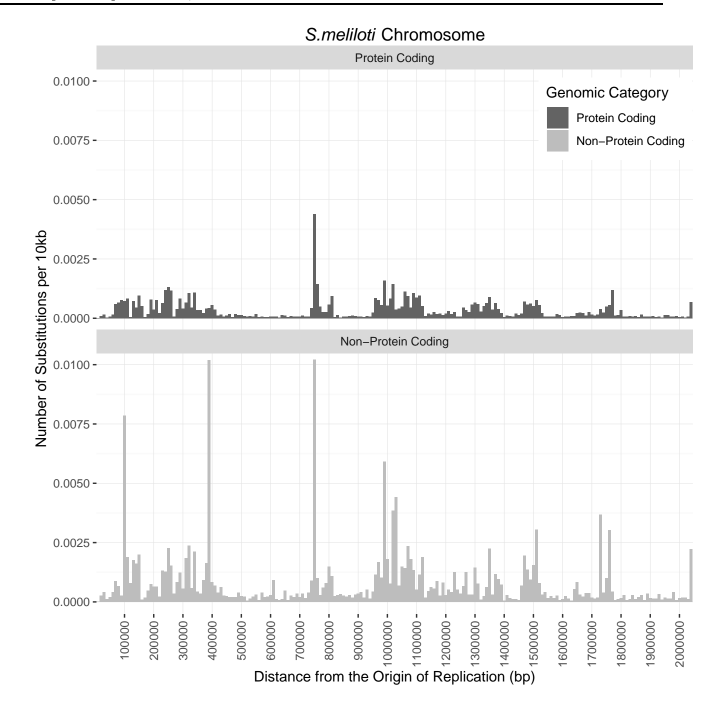
Table 1: Linear regression for dN, dS, and ω calculated for each bacterial replicon for the 20 genes closest and 20 genes farthest from the origin of replication. All results are marked with significance codes as followed: p: <0.001= '***', 0.001<0.01 = '***', 0.01<0.05 = '*', >0.05= 'NS'.

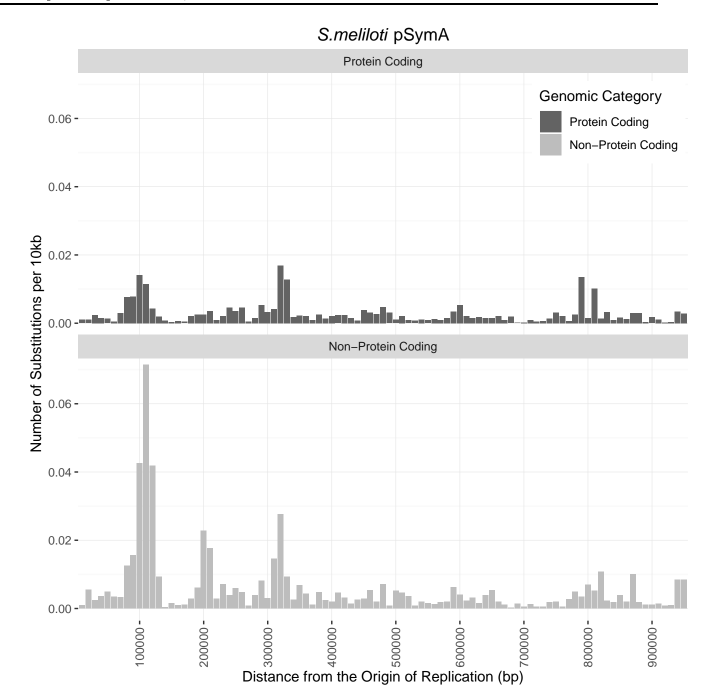


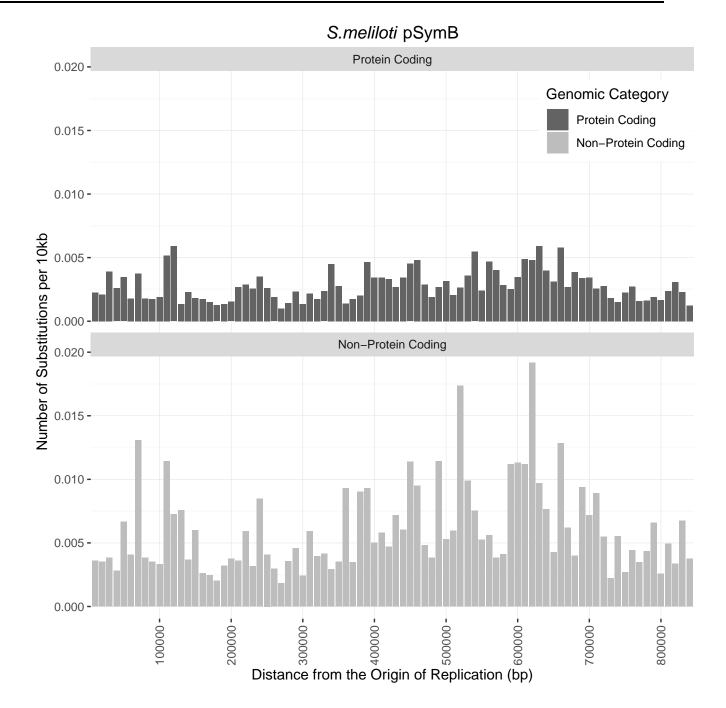


Streptomyces Chromosome









| Bacteria and Replicon | Protein Coding Sequences | Non-Protein Coding Sequences |
|--------------------------------|----------------------------|------------------------------|
| E. coli Chromosome | | NS |
| $B.\ subtilis$ Chromosome | $-4.971 \times 10^{-8***}$ | $-1.055 \times 10^{-7***}$ |
| Streptomyces Chromosome | | |
| $S.\ meliloti$ Chromosome | $-1.903 \times 10^{-7***}$ | $-2.900 \times 10^{-7***}$ |
| $S.\ meliloti\ \mathrm{pSymA}$ | $-6.642 \times 10^{-7***}$ | $-1.263 \times 10^{-6} ***$ |
| S. meliloti pSymB | $1.769 \times 10^{-7} ***$ | $4.771 \times 10^{-7} ***$ |

Table 2: Logistic regression analysis of the number of substitutions along all positions of the genome of the respective bacteria replicons. These genomic positions were split up into the coding and non-coding regions of the genome. 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 bidirectionality of replication. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "**", 0.01 < 0.05 = "*", > 0.05 = "NS".

| | Protein Coding | | | Non-Protein Coding | | | | |
|---|-----------------------------|----------------------------|--|---|--|--|---|--|
| | Correlation 20kb | | Number of Substitutions per 20kb Near | | Correlation Coefficient 20kb Near | | Number of Substitutions per 20kb Near | |
| Bacteria and Replicon | Origin | Terminus | Origin | Terminus | Origin | Terminus | Origin | Terminus |
| E. coli Chromosome B. subtilis Chromosome | $-2.889 \times 10^{-5*}$ NS | NS $1.863 \times 10^{-5*}$ | 2.87×10^{-2} | 4.24×10^{-2} | $\begin{array}{ c c c c c c c c c c c c c c c c c c c$ | $-8.209 \times 10^{-5*}$ $5.823 \times 10^{-5***}$ | $\begin{array}{ c c c c c }\hline 1.095 \times 10^{-2} \\ 8 \times 10^{-4} \\ \hline \end{array}$ | $\substack{4.45\times 10^{-3}\\6.75\times 10^{-3}}$ |
| Streptomyces Chromosome S. meliloti Chromosome S. meliloti pSymA S. meliloti pSymB | NS NS | NS NS | $\begin{vmatrix} 4.05 \times 10^{-3} \\ 6.15 \times 10^{-3} \end{vmatrix}$ | $\substack{2 \times 10^{-4} \\ 1.9 \times 10^{-3}}$ | NS 1.403×10 ^{-4***} NS | NS -2.220×10 ^{-4**} -4.557×10 ^{-5**} | $\begin{array}{c c} 9 \times 10^{-4} \\ 2.8 \times 10^{-3} \\ 5.1 \times 10^{-3} \end{array}$ | $1.5 \times 10^{-4} 5.5 \times 10^{-4} 5.4 \times 10^{-3}$ |

Table 3: Logistic regression on 20kb closest and farthest from the origin of replication after accounting for bidirectional replication and outliers. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "*", 0.001 < 0.05 = "*", 0.005 = "NS".

| | Protein | Coding | Non-Protein Coding | | |
|---------------------------------|-----------------------------|-----------------------------|----------------------------|----------------------------|--|
| Bacteria and Replicon | Weighted | Non-Weighted | Weighted | Non-Weighted | |
| E. coli Chromosome | $-4.87 \times 10^{-10} **$ | -1.839×10 ⁻⁴ *** | NS | $-2.244 \times 10^{-5***}$ | |
| B. subtilis Chromosome | NS | -2.031×10^{-4} ** | NS | $-2.885 \times 10^{-5} **$ | |
| Streptomyces Chromosome | | | | | |
| $S.\ meliloti\ { m Chromosome}$ | $-1.341 \times 10^{-10} **$ | $-1.461 \times 10^{-5} **$ | -3.490×10^{-10} * | NS | |
| $S.\ meliloti\ \mathrm{pSymA}$ | NS | NS | $-1.144 \times 10^{-8**}$ | $-6.74 \times 10^{-5} **$ | |
| S. meliloti pSymB | NS | NS | NS | NS | |

Table 4: Linear regression on 10kb sections of the genome with increasing distance from the origin of replication after accounting for bidirectional replication. Weighted columns have the total number of substitutions in each 10kb section of the genome divided by the total number of protein coding and non-protein coding sites in the genome. Non-weighted columns are performing a linear regression on the total number of substitutions in each 10kb section of the genome. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "**", 0.01 < 0.05 = "NS".

| Bacteria and Replicon | Gene Expression 10kb |
|-------------------------|----------------------------|
| E. coli Chromosome | $-2.742 \times 10^{-5**}$ |
| B. subtilis Chromosome | -2.198×10^{-5} * |
| Streptomyces Chromosome | $-5.230 \times 10^{-7***}$ |
| S. meliloti Chromosome | NS |
| S. meliloti pSymA | NS |
| S. meliloti pSymB | NS |

Table 5: Linear regression analysis of the median counts per million expression data for 10kb segments of the genome of the respective bacteria replicons. Linear 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 bidirectionality of replication. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "**", 0.01 < 0.05 = "*", > 0.05 = "NS".

| Bacteria and Replicon | Coefficient Estimate | Standard Error | P-value |
|--------------------------------|------------------------|-----------------------|------------------------------|
| E. coli Chromosome | -6.03×10^{-5} | 1.28×10^{-5} | 2.8×10^{-6} |
| B. subtilis Chromosome | -9.7×10^{-5} | 2.0×10^{-5} | 1.2×10^{-6} |
| Streptomyces Chromosome | -1.17×10^{-6} | 1.04×10^{-7} | $<2 \times 10^{-16}$ |
| $S.\ meliloti$ Chromosome | 3.97×10^{-5} | 4.25×10^{-5} | $NS (3.5 \times 10^{-1})$ |
| $S.\ meliloti\ \mathrm{pSymA}$ | 1.39×10^{-3} | 2.53×10^{-4} | 4.9×10^{-8} |
| S. meliloti pSymB | 1.46×10^{-4} | 2.03×10^{-4} | NS $(5.34.7 \times 10^{-1})$ |

Table 6: Linear regression analysis of the median counts per million expression data along the genome of the respective bacteria replicons. Linear 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 bidirectionality of replication.

| Bacteria and Replicon | Coefficient Estimate |
|-------------------------|-----------------------------|
| E. coli Chromosome | NS |
| B. subtilis Chromosome | $-2.682 \times 10^{-6} ***$ |
| Streptomyces Chromosome | $-2.360 \times 10^{-6} ***$ |
| S. meliloti Chromosome | $-2.074 \times 10^{-6} ***$ |
| S. meliloti pSymA | NS |
| S. meliloti pSymB | -4.19×10^{-6} * |

Table 7: Linear regression analysis of the total number of protein coding genes per 10kb along the genome of the respective bacteria replicons. Linear 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 bidirectionality of replication. All results are marked with significance codes as followed: < 0.001 = "**", 0.001 < 0.01 = "*", 0.01 < 0.05 = "", > 0.05 = "NS".

| Bacteria and Replicon | dN | dS | ω |
|----------------------------------|----|----|--------------------------|
| E. coli Chromosome | NS | NS | NS |
| B. subtilis Chromosome | NS | NS | -9.08×10^{-6} * |
| Streptomyces Chromosome | NS | NS | NS |
| S. meliloti Chromoeom | NS | NS | NS |
| $S.\ meliloti\ \mathrm{pSymA}$ | NS | NS | NS |
| $S. \ meliloti \ \mathrm{pSymB}$ | NS | NS | 1.163×10^{-5} * |

Table 8: Linear regression for dN, dS, and ω calculated for each bacterial replicon on a per genome basis. All results are marked with significance codes as followed: p: < 0.001 = '***', 0.001 < 0.01 < 0.01 < 0.01 < 0.05 = 'NS'.

| Bacteria and Replicon | Average Expression Value (CPM) |
|---------------------------------|--------------------------------|
| E. coli Chromosome | 160.500 |
| B. subtilis Chromosome | 176.400 |
| Streptomyces Chromosome | 6.084 |
| $S.\ meliloti$ Chromosome | 271.400 |
| $S.\ meliloti\ \mathrm{pSymA}$ | 690.100 |
| $S.\ meliloti\ p{ m Sym}{ m B}$ | 595.700 |

Table 9: Arithmetic gene expression calculated across all genes in each replicon. Expression values are represented in Counts Per Million.

| | Gene Average | | | Genome Average | | |
|----------------------------------|--------------|--------|----------|----------------|--------|----------|
| Bacteria and Replicon | dS | dN | ω | dS | dN | ω |
| E. coli Chromosome | 1.0468 | 0.1330 | 1.3183 | 0.6491 | 0.0364 | 0.2432 |
| $B.\ subtilis$ Chromosome | 4.652 | 0.2333 | 2.4200 | 1.0879 | 0.0703 | 0.3852 |
| Streptomyces Chromosome | 13.4950 | 2.0973 | 21.0423 | 5.1256 | 0.8911 | 8.9146 |
| $S.\ meliloti$ Chromosome | 0.0184 | 0.0012 | 0.1069 | 0.0187 | 0.0013 | 0.0962 |
| $S. \ meliloti \ \mathrm{pSymA}$ | 1.0602 | 0.7451 | 5.1290 | 0.4100 | 0.0863 | 0.8311 |
| $S. \ meliloti \ pSymB$ | 3.2602 | 0.0256 | 0.3878 | 0.1436 | 0.0100 | 0.1943 |

Table 10: Weighted averages calculated for each bacterial replicon on a per genome basis using the gene length as the weight. Arithmetic mean calculated for the per gene averages for each bacterial replicon.

| Bacteria Strain/Species | GEO Accession Number | Date Accessed |
|---|-----------------------|-------------------|
| E. coli K12 MG1655 | GSE60522 | December 20, 2017 |
| $E.\ coli\ \mathrm{K12}\ \mathrm{MG1655}$ | GSE73673 | December 19, 2017 |
| $E.\ coli\ \mathrm{K}12\ \mathrm{M}G1655$ | GSE85914 | December 19, 2017 |
| $E.\ coli\ \mathrm{K}12\ \mathrm{M}G1655$ | GSE40313 | November 21, 2018 |
| $E.\ coli\ \mathrm{K}12\ \mathrm{MG}1655$ | GSE114917 | November 22, 2018 |
| $E.\ coli\ \mathrm{K}12\ \mathrm{MG}1655$ | GSE54199 | November 26, 2018 |
| $E.\ coli\ \mathrm{K}12\ \mathrm{D}H10\mathrm{B}$ | GSE98890 | December 19, 2017 |
| E. coli BW25113 | GSE73673 | December 19, 2017 |
| E. coli BW25113 | GSE85914 | December 19, 2017 |
| E. coli O157:H7 | GSE46120 | August 28, 2018 |
| $E.\ coli\ { m ATCC}\ 25922$ | GSE94978 | November 23, 2018 |
| B. subtilis 168 | GSE104816 | December 14, 2017 |
| $B.\ subtilis\ 168$ | GSE67058 | December 16, 2017 |
| $B.\ subtilis\ 168$ | GSE93894 | December 15, 2017 |
| $B.\ subtilis\ 168$ | GSE80786 | November 16, 2018 |
| S. coelicolor A3 | GSE57268 | March 16, 2018 |
| $S.\ natalensis\ HW-2$ | GSE112559 | November 15, 2018 |
| S. meliloti 1021 Chromosome | GSE69880 | December 12, 2017 |
| S. meliloti 2011 pSymA | NC_020527 (Dr. Finan) | April 4, 2018 |
| S. meliloti 1021 pSymA | GSE69880 | November 15, 18 |
| S. meliloti 2011 pSymB | NC_020560 (Dr. Finan) | April 4, 2018 |
| S. meliloti 1021 pSymB | GSE69880 | November 15, 18 |
| | | |

Table 11: Summary of strains and species found for each gene expression analysis. Gene Expression Omnibus accession numbers and date accessed are provided.