Identifying Drivers of Parallel Evolution: A Regression Model Approach

Susan F. Bailey¹,²,*, Qianyun Guo¹, and Thomas Bataillon¹
¹Bioinformatics Research Centre, Aarhus University, Denmark
²Present address: Department of Biology, Clarkson University, Potsdam, NY

*Corresponding author: E-mail: sbailey@clarkson.edu.
Accepted: September 24, 2018

Abstract
Parallel evolution, defined as identical changes arising in independent populations, is often attributed to similar selective pressures favoring the fixation of identical genetic changes. However, some level of parallel evolution is also expected if mutation rates are heterogeneous across regions of the genome. Theory suggests that mutation and selection can have equal impacts on patterns of parallel evolution; however, empirical studies have yet to jointly quantify the importance of these two processes. Here, we introduce several statistical models to examine the contributions of mutation and selection heterogeneity to shaping parallel evolutionary changes at the gene-level. Using this framework, we analyze published data from forty experimentally evolved Saccharomyces cerevisiae populations. We can partition the effects of a number of genomic variables into those affecting patterns of parallel evolution via effects on the rate of arising mutations, and those affecting the retention versus loss of the arising mutations (i.e., selection). Our results suggest that gene-to-gene heterogeneity in both mutation and selection, associated with gene length, recombination rate, and number of protein domains drive parallel evolution at both synonymous and nonsynonymous sites. While there are still a number of parallel changes that are not well described, we show that allowing for heterogeneous rates of mutation and selection can provide improved predictions of the prevalence and degree of parallel evolution.

Key words: parallel evolution, experimental evolution, Poisson regression, negative binomial regression.

Introduction
Documenting patterns of parallel evolution during the adaptive divergence of populations or during repeated bouts of adaptation in populations maintained in the lab is becoming increasingly feasible. Beyond the fascination for the pattern of repeatable evolution, an outstanding open question is to understand which underlying processes are driving the pattern of molecular evolution during adaptation. Theory makes clear cut predictions: in the absence of selective interference between beneficial mutations (the so called strong selection weak mutation, or SSWM, domain), heterogeneity in mutation rates, and selection coefficients between loci are expected to have the potential for equal influence on patterns of parallel evolution (Chevin et al. 2010; Lenormand et al. 2016). In a regime where clonal interference occurs, the relative influence of heterogeneity in selection coefficients may increase, however, mutation heterogeneity still has the potential to play an important role (Bailey et al. 2017). It is known that mutation rate can vary across the genome (e.g., yeast: Lang and Murray 2008, 2011; E. coli: Lee et al. 2012), and it is very clear that selection coefficients vary across the genome, thus, we expect that in most populations both mutation and selection heterogeneity will play a role in driving patterns of parallel evolution, however, the balance of these two processes is still not clear. So far, very few empirical studies have attempted to jointly quantify the relative importance of mutation and selection in shaping patterns of parallel evolution in genetic data. One study has explored this indirectly by quantifying the contribution of these two processes in shaping the parallel evolution of heritable traits (Streiffeld and Rausher 2011). Recent work by Bailey et al. (2017) outlines an approach for quantifying the effects of mutation and selection heterogeneity in driving parallel evolution in experimental evolution data, but this alternate approach cannot identify potential genomic drivers of that heterogeneity, as we do here. Other previous studies looking explicitly at parallel genetic
Parallel evolution is an identical change in independently evolving lineages, and the similar process, convergent evolution, occurs when different ancestral states change to the same descendant state in independently evolving lineages (Zhang and Kumar 1997). These kinds of evolutionary changes are studied across many different levels of biological organization from nucleotides to genes to pathways and more. In this study, we focus on parallel evolution at the level of the gene.

Parallel, along with convergent, evolution has previously been considered strong evidence of selection. A number of studies have examined gene-level mutation counts, looking for levels of parallel evolution that exceed what one would expect in the absence of selection (Woods et al. 2006; Caballero et al. 2015; Marvig et al. 2015), according to some null model, with an aim to identify genes that are under selection. For example, Caballero et al. (2015) calculated the probability of instances of gene-level parallel evolution in whole genome sequences of Pseudomonas aeruginosa repeatedly sampled over the course of a year from the sputum of a cystic fibrosis patient assuming uniform resampling of ~150 mutation events across the ~6,000 genes in the genome. The authors were able to identify 19 different genes for which there was significant deviation from their null model, and that pattern was interpreted as evidence for selection acting on these genes. However, this study and other similar approaches do not account for the possibility of heterogeneity in mutation rate from gene-to-gene, a process that can generate false positives when using “abnormal” levels of parallel evolution as a means to detect selected genes.

Others have compared instances of parallel and convergent evolution across species (see Christin et al. 2010 for a review and examples). These studies also aim to identify genes under selection by searching for genes that exhibit a higher than expected number of instances of parallel evolution according to a specified null model for evolution. Many cross-species comparative studies report instances of parallel molecular evolution and readily interpret these as being driven by positive selection (Jost et al. 2008; Castoe et al. 2009; Liu et al. 2010; Feldman et al. 2012). However, Zou and Zhang (2015) show that in this type of analysis the choice of null model is crucial and suggest that many previously reported instances of parallel evolution driven by selection could in fact have resulted simply from mutation biases and mutational heterogeneity in the absence of selection.

In contrast to studies aimed at identifying selection, other work has focused on examining how heterogeneity in mutation rate can affect the distribution of mutations across a genome, and so the probability of parallel evolution. These studies focus exclusively on either those mutations that are assumed to be to a first approximation neutral (e.g., synonymous mutations, Maddamsetti et al. 2015) or mutations arising in the course of experiments where selection is minimal (e.g., mutations arising in a mutation accumulation experiment; Ness et al. 2015). On the whole, these studies suggest substantial gene-to-gene heterogeneity in mutation rate and this can arguably also generate differences in the distribution of mutations across the genome (although studies differ in the factors identified that drive that heterogeneity). However, it is not clear what the relative contribution of mutation rate heterogeneity is when the mutations of interest also have the potential to be under varying degrees of selection.

In this study, we aim to explore the effects of both mutation and selection in generating the mutations that are observed across the genome. By identifying and quantifying the processes that give rise to mutations and how those vary from gene-to-gene, we can then begin to understand and predict patterns of parallel evolution at the gene-level. To do this, we propose a framework that explicitly considers drivers of both selection and mutational heterogeneity. Using both Poisson and negative binomial regression models, we analyze gene-level mutation count data obtained from whole genome sequencing of a large set of yeast (Saccharomyces cerevisiae) experimental populations that were adapted in parallel to identical environmental conditions in the lab (Lang et al. 2013). We choose to work at the gene level with these data as this is more appropriate than a nucleotide level analysis given the sparsity of mutations and resolution of genomic covariates. We find that the best predictor of parallel mutations at the gene-level is simply the length of the gene, and along with this, a few other genomic covariates—namely the number of protein domains and the rate of recombination, and so it is variation in these variables that drives patterns of parallel evolution in this system.

**Models for Identifying Processes Underlying Parallel Evolution**

We are interested in quantifying heterogeneities in mutation rate and selection, and how these in turn are driving patterns of parallel evolution, and identifying genomic variables that predict how the processes of mutation and selection vary from gene-to-gene. To accomplish this, we need a framework that can explicitly separate the effects of variation in mutation rate and variation in selection. We do this by examining separately the observed synonymous and nonsynonymous mutations, making the assumption (which we then explore) that gene-to-gene variation in the rate at which synonymous mutations rise to observable frequencies is driven solely by variation in the mutation rate per gene, while gene-to-gene variation in the rate at which nonsynonymous mutations have arisen may be driven by heterogeneity in both mutation and selection processes. We describe the number of mutations observed in gene $i$ during the course of an
experiment, as \( X_i = X_{i}^{s} + X_{i}^{n} \), where \( X_{i}^{s} \) and \( X_{i}^{n} \) denote the synonymous and nonsynonymous mutation counts, respectively. We assume these mutations are Poisson distributed with rates \( \lambda_{i}^{s} \) and \( \lambda_{i}^{n} \), respectively. For synonymous mutations, this Poisson rate can be modeled as follows:

\[
\lambda_{i}^{s} = M_{0} \mu_{0} L_{i} \pi_{0}
\]

Here, \( M_{0} \) is a parameter that absorbs both time and population size at which the evolution occurred and that is constant across the genome, \( \mu_{0} \) is the per-nucleotide mutation rate that we assume (and check) is constant across the genome, \( L_{i} \) is the length of gene \( i \) in nucleotides, and \( \pi_{0} \) is the probability of a synonymous mutation rising to an observable frequency in the population (we assume that synonymous mutations are selectively neutral and so this probability is assumed to be constant across the genome). For nonsynonymous mutations,

\[
\lambda_{i}^{n} = \lambda_{i}^{s} \pi_{i},
\]

where \( \pi_{i} \) is a scalar that incorporates the effects of selection on the rate of fixation of nonsynonymous mutations arising in gene \( i \). Specifically, \( \pi_{i} \) is a function of the mean selection coefficient of gene \( i \), \( s_{i} \), and under strong-selection-weak-mutation (SSWM) conditions, \( \pi_{i} \propto s_{i} \) (Gillespie 1984). Outside of SSWM conditions, in a clonal interference regime, the reality is more like \( \pi_{i} \propto (s_{i})^{\gamma} \), where \( \gamma \geq 1 \) and increases with increased clonal interference (Bailey et al. 2017). Thus, we caution that while predictors of \( \pi_{i} \) identified using this regression approach are, indeed, predictors of variation in selection, they may be either linear or nonlinear predictors, depending on the conditions of the experiment from which data were obtained. We assume that the mean selection coefficient for nonsynonymous mutations in a given gene can range from deleterious, to neutral, to beneficial. The type of data used and the underlying assumptions are summarized in figure 1.

Given these underlying assumptions about the processes giving rise to observable mutations in the experimental sequence data, we can then use Poisson and negative binomial (NB) regression to identify potential genomic variables that significantly explain variation in \( \lambda_{i}^{n} \) and \( \lambda_{i}^{s} \), and thus ultimately in the mutation and selection processes from gene-to- gene. The Poisson regression is used to explore counts of rare events (i.e., the observed mutations) that have a fixed probability of being observed, while for a NB regression, the rate of those rare events is itself a random variable that is gamma-distributed. A NB regression incorporates an extra parameter beyond a Poisson rate, known as the dispersion parameter \( (\text{here denoted by } \theta) \), reflecting the amount of underlying variation in the rate of observed mutations from gene-to-gene and governs the “extra” variance of the NB distribution relative to a Poisson distribution with identical mean. If there is no heterogeneity among the rate of observed mutations from gene-to- gene, the dispersion parameter \( \theta \) goes to zero and

\[\text{we recover a Poisson regression model. Therefore, the Poisson regression model is a special case of the NB regression model, as NB}(\lambda_{i}, \theta) \text{ reduces to Poisson}(\lambda_{i}) \text{ at the limit of } \theta \rightarrow 0 \text{ (see for instance Zuur et al. 2009). As a consequence, the Poisson and NB models are “nested” and their relative fit can be compared using a likelihood ratio test when exploring the fit of both types of regression models in this study.}

More precisely, we use the models \( X_i \sim \text{Poisson} (\lambda_i) \) or \( X_i \sim \text{NB}(\lambda_i, \theta) \), fitting the following regression:

\[
\log (\lambda_i) = \text{constant} + x_1 A_1 + x_2 A_2 + \cdots + x_p A_p,
\]

where \( \lambda = (\lambda_1, \ldots, \lambda_n) \) are the Poisson rates for all \( n \) genes, \( A_1, \ldots, A_p \) are the \( j \) potential genomic explanatory variables, and \( x_1, \ldots, x_p \) the estimated regression coefficients for those \( j \) variables. Thus, in the case of the synonymous mutations, \( \text{constant} = \log (M_0 \mu_{0} \pi_{0}), A_1 = \log (L_i) \text{ setting } x_1 = 1 \). For nonsynonymous mutations, \( x_2 A_2 + \cdots + x_p A_p = \log (\pi_i) \).

Details of the implementation of these models is provided below.

Materials and Methods

The Data

Evolution Experiment Data

We analyzed data obtained from whole genome resequencing of forty populations of \( S. \) cerevisiae adapted in parallel in
the lab (Lang et al. 2013). In this experiment, clonal haploid yeast populations were grown in 128 μl of liquid YPD media and transferred every 12–24 h to fresh media for ~1,000 generations (see Lang et al. 2013) for more details on the experiment protocol. In our analysis, we focus on all detected genic mutations (718 out of the total 1,020 in the data set) from forty sequenced populations, that is, all genic mutations that were able to escape drift and so rise to frequencies of at least ~10% in the populations (mutations below this frequency could not reliably be detected; see Lang et al. 2013). The mutations included in the data set consisted of SNPs and small indels. Mutations were grouped by gene across all forty populations and categorized as synonymous (SYN) or nonsynonymous (NS), that is, those that do not confer amino acid changes, and those that do, respectively.

**Comparative Genomics Data**

We used a set of orthologous gene alignments spanning four distinct yeast species (S. cerevisiae, Saccharomyces paradoxus, Saccharomyces bayanus, and Saccharomyces mikatae; available from www.yeastgenome.org/download-data/genomics; Cliften 2003; Kellis et al. 2003) to infer the gene-to-gene heterogeneity of the substitution rates at synonymous sites and nonsynonymous sites, hereafter dS and dN, respectively. To do so, we first realigned the gene sequences using ClustalW (Larkin et al. 2007) on the translated protein sequence data and then applied a number of filters to the data with an aim at removing those gene alignments that might result in inaccurate codon substitution model predictions. We removed alignments for those genes where sequences were not available from all four species, alignments for which at least one sequence had <30% overlap with the one of the other three sequences, and alignments for which at least one sequence was <300 bp in length. We then used a maximum likelihood codon based method (CodeML in the PAML software package; Yang 2007) to infer dS and dN, for each gene in our data set. We used a codon table model (i.e., seqtype = 1; CodonFreq = 3) with a fixed tree topology (i.e., runmode = 0). A comparison of AICs among alternative codon-based models indicated this was the most appropriate model for the data set.

**Additional Genomic Data**

We included eight additional genomic variables in our analysis that we expected could have the potential to affect the probability of a gene to harbor mutations. Our collection of variables is not meant to be exhaustive, but simply meant to illustrate the potential for additional genomic information to improve our predictions of which genes bear mutations across the genome. For each gene, we consider: gene length, % GC content, multifunctionality, degree of protein–protein interaction (PPI), codon adaptation index (CAI), number of domains, level of expression (in the same environment at the evolution experiment), local recombination rate, and essentiality of the gene. We expect some of these variables may capture heterogeneity in the per-gene mutation rates, for example: gene length, which likely captures variation in a gene’s mutational target size, and local recombination rate, which has been shown to be associated with mutability in yeast (Strathern et al. 1995; Holbeck and Strathern 1997). We expect other variables may capture heterogeneity in selection from gene-to-gene, for example: multifunctionality and PPI, which may characterize aspects of how pleiotropic a given gene is and so the level of evolutionary constraint it is under. We expect still other genomic variables may capture heterogeneity in both mutation and selection. For example, level of expression of a gene may be correlated with gene-to-gene variation in selection as highly expressed genes have been shown to be more highly conserved, both specifically in yeast (Pál et al. 2001; Drummond et al. 2005) and as a more general phenomenon across species (Drummond and Wilke 2008). On the other hand, level of expression of a gene has also been shown to be positively correlated with mutability (Ness et al. 2015).

Descriptions of the variables used in this study and sources from which the data were obtained are provided in table 1.

For the purposes of our analysis, we only used mutations in genes for which we were able to obtain a full set of complementary genomic variables (393 of the total 718 genic mutations in Lang et al. 2013). This final data set does not contain any examples of multiple mutations within the same gene on the same genome and so we consider each mutation to be an independent mutational event. A data set integrating the mutation counts originally made available by Lang et al. (2013) (from their supplementary table S1, Supplementary Material online) and all the genomic covariates that we aggregated for this study, as well as the gene alignments used for estimating dN and dS are available on Dryad (doi: 10.5061/dryad.329b0m6).

**Regression Models**

**Regression Models and Explanatory Variables Tested**

We used the Poisson and negative binomial regression models described in the “Models” section above to examine how much of the variation in our explanatory variables could account for patterns of variation in mutation counts per gene. We used the “glm” and “glm.nb” functions in R (R Development Core Team 2014) to implement these models. We fit a series of models to synonymous and nonsynonymous mutation count data separately. To start, we fit the synonymous mutations (model M0), testing our assumptions that rate of observed mutations per gene (totaled over all 40 populations in the data set) is proportional to number of nucleotide sites in the gene (Li), and the per nucleotide mutation rate does not vary significantly across the genome—that is, a model assuming $\mu_0$ is a fixed parameter (Poisson regression) fits the data better than a model where $\mu_0$ for each gene is
Model Selection and Significance of Variables

For each variable and parameter of interest, we tested significance by comparing versions of the models with and without that variable or parameter of interest through a likelihood-ratio test (LRT). Significance testing for LRTs was done using permutation tests instead of relying on asymptotic distribution of the LRTs. Permutation was performed by resampling the mutation data, refitting the models with and without the genomic variable or parameter of interest and then calculating the LRT of those refitted models. This procedure allowed us to approximate null distributions and obtaining P values by calculating the frequency of permutations that resulted in a likelihood-ratio greater than or equal to the true observed value. Variables found to significantly improve model fit were retained in the final “best” model. We choose to test significance using permutations given that asymptotic results on the distribution of the likelihood ratio test may break down as the reduced model—the Poisson regression—lies at the boundaries of the parameter space for \( \theta \), included in the NB regression (see for instance Self and Liang 1987). In practice, 1,000 permutations were used to approximate the null and obtain P values on each variable (more permutations might be required if needed to approximate P values that are much smaller than 10\(^{-3}\)).

The two nonsynonymous mutation models \( \text{MN} \) and \( \text{MNC} \) were compared with each other using Akaike information criterion (AIC; Akaike 1973), and the proportion of variation explained (pseudo-\( R^2 \)) was estimated as the \( R^2 \) obtained from a linear regression (using “lm” in R) between the observed and predicted mutation counts for a given model. Note that this statistic is not used for any formal goodness-of-fit but as an illustrative way to report how much of the whole variation is accounted for by any model we fit to the mutation count data.
Predicting the Degree of Parallelism
We used the resulting best-fit parameters and distributions from the regression models to simulate mutations for 40 populations and calculated the mean proportion of shared genes bearing mutations for all pairwise combinations of those 40 populations using the Jaccard Index, \(J_{G1,G2} = \frac{G_{1}\cap G_2}{G_{1}\cup G_2}\). This measure of parallelism has been used in a number of previous empirical comparisons (Wong et al. 2012; Bailey et al. 2017). We compared the simulated distribution of J values to the distribution calculated from the real data set in order to assess the accuracy of the regression models in predicting the degree of parallel evolution in this system. This can be used as a predictive check for our model.

All statistical analyses were implemented in R (R Development Core Team 2014) and an example script for implementing our model framework and hypothesis testing is available on Dryad (doi:10.5061/dryad.329b0m6).

Results
The Data
Mutation Counts Data
We used experimental data comprising all mutations detected at a frequency over 10% in the forty evolved S. cerevisiae populations described in Lang et al. (2013). The mutations contained in this data set include many instances of parallel evolution at the gene level that cover a range of biological processes, the most common being negative regulation of Ras (part of a glucose signaling pathway; Santangelo 2006), cell wall biogenesis/assembly, and mating (see table 1 in Lang et al. 2013 for a comprehensive summary). After removing genes for which we had incomplete or unreliable data (see Materials and Methods), we were left with 2,891 genes out of a total of 6,603. The filtered data set contained 357 non-synonymous mutations distributed across 267 genes, and 58 synonymous mutations distributed across 57 genes. The genes removed by our filtering rules had disproportionately more mutations compared with those genes that were removed by our filtering rules had disproportionately more mutations distributed across 57 genes. This is not unexpected as highly divergent genes are more likely to be filtered out due to alignment issues, and it is not surprising that highly divergent genes would tend to see more mutations than average, whether it be as a result of mutation and/or selection mechanisms. This bias in the filtering means that our results are likely conservative in terms of detecting significant relationships between long-term (from comparative genomics data) and short-term (from experimental evolution data) measures of divergence.

Genomic Variables
We used codon substitution models comparing four yeast species to estimate dS and dN/dS for each gene. Estimates for dS ranged widely, from 0.21 to 68.7, however, the vast majority of dS estimates (~95%) were <4. Estimates for dN/dS ranged from 0.00010 to 0.43, and these values are weakly negatively correlated with dS (r = -0.043, P = 0.021). We collated and/or calculated nine other genomic variables with the potential to affect the mutation and selection processes in this system and estimated correlation coefficients between all pairs of explanatory variables used in this study (supplementary table S1, Supplementary Material online). While the correlations between these variables tend to be quite weak, many are, in fact, significant due to the large number of observations in the data set.

Mutation Counts Analysis
Synonymous Mutations
We used regression models to test our assumption that gene-level mutation rate can be adequately described as simply being directly proportional to gene length. Restricting the data to the synonymous mutations, we compared Poisson regression models with and without gene length included as an explanatory variable (\(M_{2.0.P}: \lambda_S = \text{constant}\) and \(M_{2.1.P}: \lambda_S = \text{constant} \times L(a_2, \text{respectively})\), and a Poisson regression model where rate is restricted to be directly proportional to gene length (i.e., \(M_{2.2.P}: \lambda_S = \text{constant} \times L\)). We also compared with negative binomial versions of these model to look for the possibility of additional unexplained variation in the rate \(\lambda\). A summary of the results of these comparisons is shown in table 2. Model \(M_{2.2.P}\) was the best model according to a comparison of AICs. The fits of these models to the distribution of synonymous mutation counts per gene are visualized in figure 2A. We also compared a series of Poisson models, each containing one of the genomic variables included in this study (see supplementary table S2, Supplementary Material online). Two variables do significantly improve model fit—dN/dS and CAI, but very modestly, helping to explain only 0.24% and 0.11% of the total variance, respectively. Taken together, these tests suggest that the assumptions of neutral selection at synonymous sites and a constant mutation rate across nucleotides are reasonable simplifications for these data.

Table 2

<table>
<thead>
<tr>
<th>Model</th>
<th>Log-lik.</th>
<th>No. param</th>
<th>AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>(M_{2.0.P})</td>
<td>-283.0</td>
<td>1</td>
<td>568.2</td>
</tr>
<tr>
<td>(M_{2.0.NB})</td>
<td>-283.0</td>
<td>1</td>
<td>569.9</td>
</tr>
<tr>
<td>(M_{2.1.P})</td>
<td>-273.9</td>
<td>2</td>
<td>551.8</td>
</tr>
<tr>
<td>(M_{2.1.NB})</td>
<td>-273.9</td>
<td>3</td>
<td>553.8</td>
</tr>
<tr>
<td>(M_{2.2.P})</td>
<td>-274.0</td>
<td>1</td>
<td>549.9</td>
</tr>
<tr>
<td>(M_{2.2.NB})</td>
<td>-274.0</td>
<td>2</td>
<td>551.9</td>
</tr>
</tbody>
</table>

Note.—Log-likelihoods and AIC values are provided. The best model as determined by the lowest AIC with the fewest parameters is \(M_{2.2.P}\).
We fit regression models to the nonsynonymous mutation data, including eleven genomic variables, trying to identify which of those variables could significantly explain variation in the number of observed mutations per gene (totaled over all 40 populations in the data set). We found that gene length ($L$), number of domains in the encoded protein ($num.dom$), and recombination rate ($r$) were significant in our model (see model MN.NB in table 3).

When we fit regression models using the principal components of the genomic variables in place of the raw variables, we found that only a single principal component, PC10, was significant in the model (see model MN.NBPC in table 3). PC10 is fairly evenly loaded with a number of genomic variables (see fig. 3), however, the three significant genomic variables from MN.NB ($L$, $num.dom$, and $r$) are among the variables more heavily loaded on PC10, so the two models seem to be roughly in agreement. Further, these models both explain $\sim$16% of the variance (as calculated from pseudo-$R^2$ estimates, see Materials and Methods). A comparison of Poisson and negative binomial regression models, as well as models including the raw genomic variables versus the transformed principal component variables, suggests that the best model for these nonsynonymous mutation count data is a negative binomial regression using the raw genomic variables (see AIC values in table 4). The fits of these models to the

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**Table 3**

“MN” Models Parameter Estimates (constant, $a_1$, $a_2$, etc.) and $P$-values for Those Estimates

<table>
<thead>
<tr>
<th>Model</th>
<th>Parameter</th>
<th>Estimate</th>
<th>$P$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MN.NB</td>
<td>$\lambda_N = \text{constant} \times L_i \times L_i^n \times \text{num.dom}_i^n \times r_i^n \times \theta_h$</td>
<td>$a_1 = 0.4405$</td>
<td>$0.001$</td>
</tr>
<tr>
<td></td>
<td>$\lambda_N \sim L$</td>
<td>$a_2 = -0.4511$</td>
<td>$0.004$</td>
</tr>
<tr>
<td></td>
<td>$\lambda_N \sim num.dom$</td>
<td>$a_3 = 0.1029$</td>
<td>$0.041$</td>
</tr>
<tr>
<td></td>
<td>$\lambda_N \sim r$</td>
<td>$a_3 = 0.1029$</td>
<td>$0.041$</td>
</tr>
<tr>
<td></td>
<td>$\lambda_N \sim \text{constant}$</td>
<td>$8.084 \times 10^{-6}$</td>
<td>$&lt; 0.001$</td>
</tr>
<tr>
<td></td>
<td>$\lambda_N \sim \delta_N$</td>
<td>$0.3806$</td>
<td>$&lt; 0.001$</td>
</tr>
</tbody>
</table>

| MN.NBPC | $\lambda_N = \text{constant} \times L_i \times \exp(\text{PC10}_i^{\beta_1}) \times \theta_h$ | $\exp(\text{PC10}) = 0.2984$ | $< 0.001$ |
|         | $\lambda_N \sim \exp(\text{PC10})$ | $\beta_1 = 0.2984$ | $< 0.001$ |
|         | $\lambda_N \sim \text{constant}$ | $8.846 \times 10^{-5}$ | $< 0.001$ |
|         | $\lambda_N \sim \delta_N$ | $0.3988$ | $< 0.001$ |

*NOTE.*—Only those variables that significantly improved model fit are included.

**Nonsynonymous Mutations**

We fit regression models to the nonsynonymous mutation data, including eleven genomic variables, trying to identify which of those variables could significantly explain variation in the number of observed mutations per gene (totaled over all 40 populations in the data set). We found that gene length ($L$), number of domains in the encoded protein ($num.dom$), and recombination rate ($r$) were significant in our model (see model MN.NB in table 3).

When we fit regression models using the principal components of the genomic variables in place of the raw variables, we found that only a single principal component, PC10, was significant in the model (see model MN.NBPC in table 3). PC10 is fairly evenly loaded with a number of genomic variables (see fig. 3), however, the three significant genomic variables from MN.NB ($L$, $num.dom$, and $r$) are among the variables more heavily loaded on PC10, so the two models seem to be roughly in agreement. Further, these models both explain $\sim$16% of the variance (as calculated from pseudo-$R^2$ estimates, see Materials and Methods). A comparison of Poisson and negative binomial regression models, as well as models including the raw genomic variables versus the transformed principal component variables, suggests that the best model for these nonsynonymous mutation count data is a negative binomial regression using the raw genomic variables (see AIC values in table 4). The fits of these models to the
distribution of nonsynonymous mutation counts per gene are visualized in figure 2B.

Predicted Parallelism

Using the predicted rates and distributions from the best-fit regression models for both the synonymous and nonsynonymous mutation counts (M₃,2.P and M₃,4.NB, respectively), we simulated mutations for a set of 40 populations and calculated the Jaccard index (J) as a measure of gene-level parallelism between all pairs of those simulated populations. We performed the same calculations with the 40 populations from the real data. Figure 4 shows a comparison of those J values from the simulated and real data. While the simulated data from our model does quite well at capturing the range of J values, it does not do a great job of capturing the shape of the distribution.

Discussion

Here, we present a modeling framework to infer what genomic variables may underlie gene-to-gene variation in mutation rate and intensity of selection. We use these models to provide evidence that parallel evolution at both nonsynonymous and synonymous sites is driven by nontrivial amounts of gene-to-gene heterogeneity in mutation, selection, and the combination of the two processes. Using our modeling approach, we identified a number of genomic variables and their potential contributions to the probability of parallel evolution via the processes of mutation and selection.

Longer Genes Harbor More Mutations

By far, the variable having the largest effect on variation in the number of synonymous and nonsynonymous mutations observed was gene length. More specifically, gene length positively affected the rate of mutation at the gene-level, meaning genes comprising more nucleotides were more likely to harbor mutations. This result is not surprising and is in agreement with analysis of synonymous mutation counts from Lentzki’s long-term evolution experiment with E. coli (Maddamsetti et al. 2015).

Long-Term Divergence Does Not Predict Short-Term Mutation Counts

Our model for synonymous mutation counts suggests that divergence estimates from long-term evolutionary comparisons at the species level do not provide insight into expected mutation counts on the shorter time scale of evolution in the lab, also in agreement with recent analysis of E. coli data (Maddamsetti et al. 2015). Maddamsetti et al. found that their proxy for long-term per gene mutation rate, θₛ (a measure of within-species nucleotide diversity), did not explain gene-to-gene variation in synonymous mutation counts in their data. The authors argued that horizontal gene transfer (HGT) is therefore likely a more important process driving gene-to-gene variation in long-term divergence between naturally occurring E. coli strains, and since HGT did not occur in their evolution experiment, it is not surprising that the experiment’s synonymous mutation counts did not correlate with θₛ. However, rates of HGT tend to be higher in bacteria, and in particular E. coli, as compared with yeast and other eukaryotes (Boto 2010). Furthermore, a recent mutation accumulation experiment with the eukaryote Chlamydomonas reinhardtii showed a positive correlation between a proxy for long-term mutation rate (θₛ) and per site mutability (Ness et al. 2015). Thus, it is somewhat surprising that we do not see a significant relationship between dS and dNdS and counts of synonymous and nonsynonymous mutations, respectively, in our examination of the S. cerevisiae data used in this study. One possibility might also be that dS and dNdS are noisy to estimate at the gene level and that tends to downplay their predictive power in our analysis of counts in an evolve and resequence experiment.

Table 4

<table>
<thead>
<tr>
<th>Model</th>
<th>Log-lik</th>
<th>No. param.</th>
<th>AIC</th>
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<tr>
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<td>2,321.7</td>
</tr>
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<td>4</td>
<td>2,050.8</td>
</tr>
<tr>
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<tr>
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<tr>
<td>M₃,NBPC</td>
<td>-947.9</td>
<td>3</td>
<td>1,901.8</td>
</tr>
</tbody>
</table>

Note.—The best model as determined by the lowest AIC with the fewest parameters is M₃,NB.
Nonsynonymous Mutation Counts Show Evidence of Selection Heterogeneity

As expected (Lenormand et al. 2016), we report strong evidence that the distribution of nonsynonymous mutations across the genome was driven in part by gene-to-gene heterogeneity in selection. Of those genomic variables tested, we found three that were significant predictors of nonsynonymous mutation counts, suggesting that those variables may drive or are correlated with processes that modulate the intensity of selection across genes. The significant variables were gene length, recombination rate, and number of protein domains.

We found that gene length predicts nonsynonymous mutation count via selection, over and above its effects on per gene mutation rate—as estimated from models aimed at explaining the synonymous mutation count only. While one might not expect gene length to have direct effects on selection, we suggest that gene length may show a significant effect here because it is correlated with other attributes of the genome that could have important effects on selection, for example, gene expression levels and multifunctionality. Because of these correlations, it could be that gene length acts as a kind of summary variable for these covariates and other unidentified factors we have not captured in these models. In fact, it is almost certainly the case that to some degree all the significant variables in our model summarize variation from additional unknown factors we have not included in our data set.

In contrast to the positive relationship between gene length and number of nonsynonymous mutations, we also found that the number of protein domains for a gene codes for (a variable that is positively correlated with gene length; supplementary table S1, Supplementary Material online) actually negatively predicts the number of nonsynonymous mutations. In other words, the more domains in the encoded protein of a gene, the fewer mutations that gene is expected to incur in the course of the yeast evolution experiment analyzed here. The mechanism behind this effect is not clear, but certainly protein structure has previously been reported to have significant impacts on evolutionary rates in yeast (Bloom et al. 2006) and one can also posit that genes encoding proteins with multiple domains and thereby involved in more numerous interactions are—all else being equal—more severely constrained by purifying selection. It is interesting that this effect can be observed in the course of relatively short time span (relative to between species divergence times) through the relative paucity of nonsynonymous mutations in these genes.

Our analysis also showed that recombination rate is a significant predictor of the observed number of nonsynonymous mutations observed in a given gene in these data. Genes with higher recombination rates are more likely to bear nonsynonymous mutations. We expect recombination rate to be correlated with mutation, as previous studies in yeast have shown that recombinational repair of double strand breaks substantially increases the frequency of point mutations in nearby intervals (Strathern et al. 1995; Holbeck and Strathern 1997). However, it is not clear why regions of the genome with high recombination rates would be correlated with selection in this data set. The recombination rate data used in our analysis comes from work that inferred recombination using crosses of natural yeast populations (Illingworth et al. 2013). The experimental yeast populations in this data set where propagated asexually so recombination is unlikely to be directly impacting evolution in this system, but instead variation in recombination rate must be associated with other processes that drive selection in this data set. Another non-exclusive possibility might be the fact that biased gene conversion might vary from gene to gene and also—like selection—augment the probability of detecting variants in evolve and resequence experiments. However, if this was the case, we might expect GC content to be a significant
predictor of both synonymous and nonsynonymous mutations in our models, and it is not ($S: P = 0.125, NS: P = 0.221$). We might also expect to see a bias toward GC in the observed mutations, however, this was not the case—only 33% of the SNPs in this data set were a change to G or C.

**Factors Driving Mutation and Selection Are Complex**

It is difficult to obtain any additional insights from models that include principal components of the genomic covariate data, however, there is at least some level of agreement between those variables that are significant (i.e., length, recombination, and number of domains) and ones that are heavily weighted in PC10—the principal component that was found to be significant (see fig. 3). The local properties of the genome do appear to drive some heterogeneity in the selection processes, and in turn, shape the patterns of parallel evolution, however, individual effects that can be ascribed to individual variables are not easy to parse out.

We want to stress that while we were able to identify a number of factors affecting the count of mutations observed in this evolution experiment data set, the total explained variance is still low: 1% and 16.0% in the synonymous and nonsynonymous models, respectively (calculated from pseudo-$R^2$ estimates of the “best” models, see Materials and Methods). While the models do capture the general distribution of mutation counts (fig. 2) and so the degree of parallel evolution, accurately predicting on which genes those mutations will fall is still very difficult. This is not surprising given the amount of stochasticity involved in both the origin of new mutations and their evolutionary fate through drift and selection. A clearer picture might emerge when using our modeling approach in a meta-analysis approach where several evolve and resequence experiments are considered together (see Bailey et al. 2017 for a similar approach on summary statistics of the amount of parallel evolution at the gene level across a wide range of experimental studies in yeast and bacteria).

While we do find a number of genomic variables that significantly affect the distribution of mutations across the genome, it is noteworthy that these models are still unable to capture the more extreme patterns of parallel evolution observed in this data set. For example, one gene (IRA1) saw mutations in over 50% of the populations sequenced in this experimental data set (discussed in more detail in Lang et al. 2013). Such a mutation count is completely out of the range of likely outcomes predicted by our models. Some of this discrepancy may be because of the simplifying assumptions made about the process of selection. Our framework models the process of mutation and its heterogeneity but while we account for the fact that newly arising mutations may have different probabilities of reaching an observable frequency, the modeling of that process could be made more precise by incorporating an explicit underlying distribution of fitness effects of new mutations at each gene. Incorporating a selection process that allows for different amounts of both positive and negative selection, as well as further details about the selection pressures in the particular environment of interest—something we do not consider at all in this study—would likely improve prediction for some of these more extreme events.

It is also important to note that the methods used in this study are focused on parallelism in SNPs and small indels. While this focus is appropriate for the data set used here, it may not be appropriate for other systems. For example, in an experimental evolution study with E. coli, (Tenaillon et al. 2012) saw that much of the parallelism seen between populations was the result of IS elements and large scale duplication and deletions. It may be important to try to account for this more diverse range of mutational event types when trying to identify the drivers of parallel evolution other systems.

Can we use this modeling framework to predict parallel evolution? To some degree, yes—the measures of parallel evolution between populations simulated using our model predictions span a similar range to those calculated from the real population data (see fig. 4). However, while this congruence suggests, we are on the right track, the shapes of the real and simulated distributions are still quite different. For example, there is, on an average, more parallelism between the real populations compared with populations simulated from our models, and in particular, there seems to be a substantial discrepancy between the number of real population-pairs and simulated population-pairs that have a low level of parallelism (e.g., note the difference between real and simulated when $J$ ranges from 0.025 to 0.05 in fig. 4B). This is further evidence suggesting that although our current models may be useful to some extent, we are still missing some important factors driving heterogeneity in mutation and selection across these genomes.

**Advantages of This Regression Framework**

Relying on the assumption that synonymous mutations are selectively neutral (which does appear to be appropriate for these data), the regression models we use in this study allow us to distinguish between genomic variables influencing the observed distribution of mutations across a genome through their potential effects on both gene-to-gene heterogeneity in mutation rate and gene-to-gene heterogeneity in selection. The great advantage of this is that it allows us to begin to break down the importance of these two processes in shaping patterns of parallel evolution we see and move closer to the goal of predicting which genes will be involved in evolution when organisms adapt to new environments. It will be interesting to apply this model framework to other data sets of this type, as they become available, to see how general these patterns are across different organisms and selection environments (Bailey and Bataillon 2016).
Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

Acknowledgments

We thank Stéphanie Bedhomme, Bastien Bousso, and three anonymous reviewers for their constructive comments on earlier versions of this manuscript. This work was supported by the European Research Council under the European Union’s Seventh Framework Program (FP7/20072013, ERC grant number 311341 to T.B.).

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Associate editor: Ruth Hershberg